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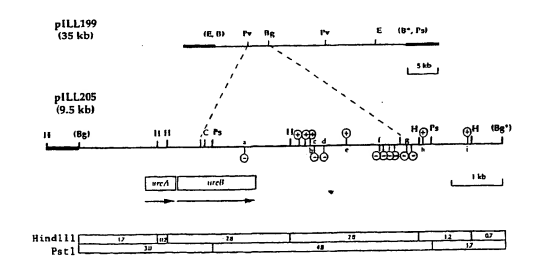
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(57) Abstract

The invention relates to an immunogenic composition, capable of inducing protective antibodies against Helicobacter infection, characterised in that it comprises: i) at least one sub-unit of a urease structural polypeptide from Helicobacter pylori, or a fragment thereof, said fragment being recognised by antibodies reacting with Helicobacter felis urease, and/or at least one sub-unit of a urease structural polypeptide from Helicobacter felis, or a fragment thereof, said fragment being recognised by antibodies reacting with Helicobacter pylori urease; ii) and/or, a Heat Shock protein (HSP), or chaperonin, from Helicobacter, or a fragment of said protein. The invention also relates to the preparation, by recombinant means, of such immunogenic compositions.

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IMMUNOGENIC COMPOSITIONS AGAINST HELICOBACTER INFECTION, POLYPEPTIDES FOR USE IN THE COMPOSITIONS AND NUCLEIC ACID SEQUENCES ENCODING SAID POLYPEPTIDES

present invention relates to immunogenic compositions for inducing protective antibodies against Helicobacter spp. infection. It also relates to proteinaceous material derived from Helicobacter, nucleic acid sequences encoding them. Antibodies to these proteinaceous materials are also included in the invention.

H. pylori is a microorganism which infects human gastric mucosa and is associated with active chronic gastritis. It has been shown to be an aetiological agent in gastroducdenal ulceration (Peterson, 1991) and two recent studies have reported that persons infected with H. pylori had a higher risk of developing gastric cancer (Nomura et al, 1991; Parsonnet et al, 1991).

In vivo studies of the bacterium and, consequently, work on the development of appropriate preventive or therapeutic agents has been severely hindered by the fact that <u>Helicobacter pylori</u> only associates with gastric-type epithelium from very few animal hosts, none of which are suitable for use as laboratory models.

A mouse model of gastric colonisation has been developed using a helical bacterium isolated from cat gastric mucus (<u>Lee et al</u>, 1988, 1990) and identified as a member of the genus <u>Helicobacter</u>. It has been named <u>H. felis</u> (<u>Paster et al</u>, 1990).

To date, only limited information concerning <u>H.</u> <u>felis</u> and the extent of its similarities and

differences with <u>H. pylori</u>, is available. The reliability of the mouse model for the development of treatments for <u>H. pylori</u> infection is therefore uncertain. Recently, it was shown that <u>H. pylori</u> urease is a protective antigen in the <u>H. felis</u> / mouse model (<u>Davin et al</u>, 1993).

It is therefore an aim of the present invention to provide therapeutic and preventive compositions for use in <u>Helicobacter</u> infection, which furthermore can be tested in laboratory animals.

It is known that <u>H. pylori</u> expresses urease activity and that urease plays an important role in bacterial colonisation and mediation of certain pathogenic processes (<u>Ferrero and Lee</u>, 1991; <u>Hazel et</u> al, 1991).

The genes coding for the urease structural polypeptides of <u>H. pylori</u> (<u>URE A, URE B</u>) have been cloned and sequenced (<u>Labigne et al</u>, 1991; and French Patent Application FR 8813135), as have the genes coding the "accessory" polypeptides necessary for urease activity in <u>H. pylori</u> (International patent application WO 93/07273).

Attempts have been made to use nucleic acid sequences from the <u>H. pylori</u> urease gene cluster as probes to identify urease sequences in <u>H. felis</u>. However, none of these attempts have been successful. Furthermore, the establishment and maintenance of <u>H. felis</u> cultures <u>in vitro</u> is extremely difficult, and the large quantities of nucleases present in the bacteria complicates the extraction of DNA.

The present inventors have however, succeeded in cloning and sequencing the genes of the urease structural polypeptides of <u>H. felis</u>, and of the accessory polypeptides. This has enabled, in the

context of the invention, the comparison of the amino-acid sequence data for the <u>H. felis ure</u> gene products with that for <u>Helicobacter pylori</u>, and a high degree of conservation between the urease sub-units has been found. An immunological relationship between the 2 ureases exists, and protective antibodies to <u>Helicobacter</u> infection can be induced using the urease sub-units or fragments thereof as immunogens.

Indeed, to elucidate the efficiency of individual urease subunits to act as mucosal immunogens, genes encoding the respective urease subunits (UreA and UreB) of Helicobacter pylori and Helicobacter felis have been cloned in an expression vector (pMAL), and expressed in Escherichia coli cells translational fusion proteins. The recombinant UreA and UreB proteins have been purified by affinity and anion exchange chromatography techniques, and have predicted molecular weights of approximately 68 and 103 kDa, respectively. Western blotting indicated that the urease components of the fusion proteins are strongly immunogenic and are specifically recognized by polyclonal rabbit anti-Helicobacter sera. Orogastric immunization of mice with 50 μg of recombinant H. felis UreB, administered combination with a mucosal adjuvant (cholera toxin), protected 60 % (n = 7; p < 0.005) of gastric colonization by \underline{H} . \underline{felis} bacteria at over 4 months. This compared with a value of 25 % (n = 8; p > 0.05) for the heterologous <u>H. pylori</u> UreB antigen. For the first time, a recombinant subunit antigen has been shown to induce an immunoprotective response against gastric Helicobacter infection.

The inventors have also identified, in the context of the invention, new Heat Shock Proteins or chaperonins, in <u>Helicobacter</u>, which have an enhancing

effect on urease activity. Use of the chaperonins in an immunogenic composition may induce therefore an enhancement of protection.

Indeed, the genes encoding each of the HspA and HspB polypeptides of Helicobacter pylori have been cloned, expressed independently as fused proteins to the Maltose-Binding-Protein (MBP), and purified on a These proteins have been used large scale. recombinant antigens to immunize rabbits, and Western immunoblotting assays as well as ELISA to determine their immunogenicity in patients infected with HP (HP+). The MBP-HspA and MBP-HspB fusion proteins have been shown to retain their antigenic properties. Comparison of the humoral immune response against HspA and/or HspB in (HP+) patient sera demonstrated that not only HspB but also HspA was recognized by (HP+) patient sera (29/38 and 15/38, respectively). None of the 14 uninfected patients had antibodies reacting with the Hsps.

The present invention concerns an immunogenic composition capable of inducing antibodies against Helicobacter infection characterised in that it comprises:

- i) at least one sub-unit of a urease structural polypeptide from <u>Helicobacter pylori</u>, or a fragment thereof, said fragment being recognised by antibodies reacting with <u>Helicobacter felis</u> urease, and/or at least one sub-unit of a urease structural polypeptide from <u>Helicobacter felis</u>, or a fragment thereof, said fragment being recognised by antibodies reacting with <u>Helicobacter pylori</u> urease;
- ii) and/or a Heat Shock protein (HSP), or chaperonin, from <u>Helicobacter</u>, or a fragment of said protein.

Preferably, the immunogenic composition is capable of inducing protective antibodies.

preferred embodiment, to According a immunogenic composition of the invention contains, as the major active ingredient, at least one sub-unit of urease structural polypeptide from Helicobacter pylori and/or Helicobacter felis. The expression "urease structural polypeptide" signifies, in the context of the present invention, enzyme of the Helicobacter pylori or Helicobacter felis probably a major surface antigen composed of two monomeric sub-units, a major sub-unit (product of the ure B gene) and a minor sub-unit, product of the ure A gene and which, when complemented by the presence of the products of the accessory genes of the urease gene cluster, are responsible for urease activity i.e. the in the two hydrolysis of urea to liberate NH4* Helicobacter species. It is to be understood that in the absence of the accessory gene products, the urease structural polypeptides do not exhibit activity, but are recognised by antibodies reacting with H. felis or H. pylori urease.

The term "immunogenic composition" signifies, in the context of the invention, a composition comprising a major active ingredient as defined above, together with any necessary ingredients to ensure or to optimise an immunogenic response, for example adjuvants, such as mucosal adjuvant, etc...

structural Helicobacter pylori urease The polypeptide has been described and sequenced by Labigne et al, 1991. The polypeptide described in this paper is particularly appropriate for use in the invention. However, the present composition of functional homology showing variants published sequence may be used, which comprise aminoacid substitutions, deletions or insertions provided that the immunological characteristics of the polypeptide in so far as its cross-reactivity with anti-Helicobacter felis urease antibodies is concerned, are maintained. Generally speaking, the polypeptide variant will show a homology of at least 75% and preferably about 90% with the included sequence.

A fragment of the <u>Helicobacter pylori</u> urease structural polypeptide may also be used in the immunogenic composition of the invention, provided that the fragments are recognised by antibodies reacting with <u>Helicobacter felis</u> urease. Such a fragment will generally be comprised of at least 6 amino-acids, for example, from 6 to 100 amino-acids, preferably about 20-25. Advantageously, the fragment carries epitopes unique to <u>Helicobacter</u>.

Nucleic acid and amino-acid sequences may be interpreted in the context of the present invention by reference to figures 11 and 12, showing the genetic code and amino-acid abbreviations respectively.

Helicobacter felis urease polypeptide suitable for use in the present invention is preferably that encoded by part of the plasmid pILL205 (deposited at the CNCM on 25th August 1993, under number: CNCM I-1355), and whose amino-acid sequence is shown in figure 3 (subunits A and B). Again, a variant of this polypeptide comprising amino-acid substitutions, deletions or insertions with respect to the figure 3 sequence may be used provided immunological cross-relationship the that Helicobacter pylori urease is maintained. Such variant normally exhibits at least 90 % homology or identity with the figure 3 sequence. An example of such variants are the urease A and B sub-units from Helicobacter heilmannii (Solnick et al, 1994), shown to have 80 % and 92 % identity with the H. felis urease A and B sub-units, respectively.

Fragments of this urease or variants may be used in the immunogenic composition provided that the fragments are recognised by antibodies reacting with Helicobacter pylori urease. Again, the length of such a fragment is usually at least 6 amino-acids, for example from 6 to 100, preferably about 20 to 25. Preferably, the fragment carries epitopes unique to Helicobacter.

If variants or fragments of the native urease sequences are employed in the immunogenic composition the invention, their cross-reactivity reacting with urease from the other antibodies Helicobacter species can be tested by contacting the fragment or the variant with antibodies, preferably polyclonal raised to either the native the recombinant urease or, alternatively, Helicobacter. Preferably, the variants and fragments give rise to antibodies which are also capable of reacting with H. heilmannii urease. Cross protection to infection by H. heilmannii is therefore also obtained by the immunogenic composition of the invention.

The use of fragments of the urease structural genes is particularly preferred since the immunological properties of the whole polypeptide may be conserved whilst minimizing risk of toxicity.

The active component of the immunogenic composition of the invention may be comprised of one sub-unit only of the urease structural polypeptide, that is either sub-unit A or sub-unit B products of the ure A and ure B genes respectively. Compositions comprising only the urease sub-unit Ure B, of either

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H. pylori or H. felis, or variants and fragments as defined above, are particularly advantageous. Most preferred are homologous systems wherein the urease sub-unit particularly sub-unit B, is derived from the organism against which protection is sought, e.g. H. felis sub-unit B against H. felis infection. However, the composition may contain both A and B sub-units, which are normally present as distinct polypeptides. However, it is possible, when the polypeptide is produced by recombinant means, to use a fusion protein comprising the entire sequences of the A and B gene products by the suppression of the stop-codon separating the two adjacent coding sequences.

component of the immunogenic urease composition, whether sub-unit A or sub-unit B, may be used in the form of translational fusion proteins, for example with the Maltose-Binding-Protein (MBP). Other suitable fusions are exemplified in International Patent Application WO 90/11360. Another example of a suitable fusion protein is the "QIAexpress" system commercialised by QIAGEN, USA, which allows the 6xHis tag sequence to be placed at the 5' or 3' end of the protein coding sequence. The use of the ingredients in the form of fusion proteins is however, entirely optional.

According to a further preferred embodiment, the immunogenic composition of the invention may comprise in addition to or instead of the urease structural polypeptide defined above, a Heat Shock Protein also known as a "chaperonin" from <u>Helicobacter</u>. These chaperonins have been elucidated by the inventors in the context of the present invention. Preferably, the chaperonin is from <u>Helicobacter pylori</u>. Such an HSP may be the urease-associated HSP A or HSP B or a mixture of the two, having the amino-acid sequence

illustrated in figure 6. These polypeptides are encoded by the plasmid pILL689 (deposited at CNCM on 25th August 1993, under number: CNCM I-1356). Particularly preferred is the <u>H. pylori</u> HSP-A protein, either alone or in combination with Hsp-B.

It is also possible to use, as HSP component, according to the invention, a polypeptide variant in which amino-acids of the figure 6 sequence have been replaced, inserted or deleted, the said variant normally exhibiting at least 75 %, and preferably at least 85 % homology with the native HSP. The variants preferably exhibit at least 75 %, for example at least 85 % identity with the native Hsp.

The variants may further exhibit functional homology with the native polypeptide. In the case of the HSP components, "functional homology" means the capacity to enhance urease activity in a organism capable of expressing active urease, and/or the capacity to block infection by Helicobacter, particularly H. felis and H. pylori. The property of enhancing urease activity may be tested using the quantitative urease activity assay described below in the examples. Fragments of either or both of the HSP A and HSP B polypeptides preferably having at least 6 amino-acids, may be used in the composition. fragments or variants of the HSP component used in the immunogenic composition of the invention are preferably capable of generating antibodies which block the urease enhancing effect normally exhibited by the HSPs. This property is also tested using the quantitative assay described in the examples. the chaperonins presence of in the composition enhances the protection against Helicobacter pylori and felis.

The Hsp component of the immunogenic composition, whether HspA or HspB can be used in the form of a translational fusion protein, for example with the Maltose-Binding-Protein (MBP). As for the suitable fusion partners are component, other WO International Patent Application described in 90/11360. The "QIAexpress" system of QIAGEN, USA, may also be used. Again, the use of the proteins in the form of fusion proteins is entirely optional.

According to the invention therefore the immunogenic composition may comprise either a urease structural polypeptide as defined above, or a <u>Helicobacter</u> Hsp, particularly HspA or a combination of these immunogens.

preferred embodiment, the According to a composition comprises, as immunogenic the A and B sub-units of both component, both Helicobacter felis (i.e. without H. pylori urease) together with the HSP A and HSP B of Helicobacter pylori. Alternatively, the A and B sub-units of the Helicobacter felis urease may be used together with those of H. pylori, but without chaperonin component.

The immunological cross-reactivity between the ureases of the two different <u>Helicobacter</u> species enables the use of one urease only in the composition, preferably that of <u>Helicobacter felis</u>. The protective antibodies induced by the common epitopes will however be active against both <u>Helicobacter pylori</u> and <u>Helicobacter felis</u>. It is also possible that the composition induce protective antibodies to other species of <u>Helicobacter</u>, if the urease polypeptide or fragment carries epitopes occuring also on those other species.

The composition of the invention is advantageously used as an immunogenic composition or a

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vaccine, together with physiologicaly acceptable excipients and carriers and, optionally, with adjuvants, haptens, carriers, stabilizers, etc. Suitable adjuvants include muranmyl dipeptide (MDP), complete and incomplete Freund's adjuvants (CFA and IFA) and alum. The vaccine compositions are normally formulated for oral administration.

The vaccines are preferably for use in man, but may also be administered in non-human animals, for example for vetinary purposes, or for use in laboratory animals such as mice, cats and dogs.

The immunogenic compositions injected into animals raises the synthesis in vivo of specific antibodies, which can be used for therapeutic purposes, for example in passive immunity.

The invention also relates to the proteinaceous materials used in the immunogenic composition and to proteinaceous material encoded by the urease gene clusters other than the A and B urease structural sub-units. "Proteinaceous material" means any molecule comprised of chains of amino-acids, eg. peptides, polypeptides or proteins, fusion or mixed proteins (i.e. an association of 2 or more proteinaceous materials, all or some of which may have immunogenic or immunomodulation properties), either purified or in mixture with other proteinaceous or proteinaceous material. "Polypeptide" signifies chain of amino-acids whatever its length and englobes the term "peptide". The term "fragment" means any amino-acid sequence shorter by at least one amino-acid than the parent sequence and comprising a length of amino-acids e.g. at least 6 residues, consecutive in the parent sequence.

The peptide sequences of the invention, may for example, be obtained by chemical synthesis, using a

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technique such as the Merrifield technique and synthesiser of the type commercialised by Applied Biosystems.

In particular, the invention relates to proteinaceous material characterised in comprises at least one of the Helicobacter felis polypeptides encoded by the urease gene cluster of the plasmid pILL205 (CNCM I-1355), including structural and accessory urease polypeptides, or a polypeptide having at least 90 % homology with said polypeptides, or a fragment thereof. Of particular interest are the gene products of the $\underline{ure \ A}$ and $\underline{ure \ B}$ as illustrated in figure 3, or a variant thereof having at least 90 % homology or a fragment having at least 6 amino-acids. The fragments and the variants are recognised by antibodies reacting with Helicobacter pylori urease.

Amongst the polypeptides encoded by the accessory genes of the urease gene cluster, is the gene product of ure I, as illustrated in figure 9, which also forms part of the invention. Also included is a variant of the ure I product having at least 75 % homology, preferably at least 85 %, or a fragment of the gene product or of the variant having at least 6 aminoacids. The variant preferably has the capacity to activate the ure A and ure, B gene products in the of the remaining urease accessory presence products. This functional homology can be detected by using the following test : 109 bacteria containing the ure I gene product variant are suspended in 1 ml of urea-indole medium and incubated at 37° C. hydrolysis of the urea leads to the release of ammonium, which increases pH and induces a colour change from orange to fuscia-red. The observation of such a colour change demonstrates that the variant of

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the \underline{ure} I gene product under test is capable of activating the \underline{ure} A and B gene products.

It is also possible that a fragment of the <u>ure I</u> gene product, if it has a length of, for example, at least 70 or 100 amino-acids, may also exhibit this functional homology with the entire polypeptide.

The fragments of <u>ure I</u> polypeptide or of the variant preferably are capable of inducing the formation of antibodies which block the urease maturation process. In other words, the fragments bear epitopes which play a decisive role in the interaction between the <u>ure I</u> and <u>ure A / ure B</u> gene products.

The invention also relates to the proteinaceous material comprising at least one of the Heat Shock Proteins or chaperonins of <u>Helicobacter pylori</u> or a fragment thereof. Particularly preferred are the HSP A and HSP B polypeptides as illustrated in figure 6 or a polypeptide having at least 75 %, and preferably at least 80 or 90 %, homology or identity with the said polypeptide. A particularly preferred fragment of the <u>Helicobacter pylori</u> HSP A polypeptide is the C-terminal sequence:

G S C C H T G N H D H K H A K E H E A C C H D H K K H

or a sub-fragment of this sequence having at least 6 consecutive amino-acids. This C-terminal sequence is thought to act as a metal binding domain allowing binding of, for example, nickel.

The proteinaceous material of the invention may also comprise or consist of a fusion or mixed protein including at least one of the sub-units of the urease structural polypeptide of \underline{H} . \underline{pylori} and/or of \underline{H} . \underline{felis} , or fragments or variants thereof as defined above. Particularly preferred fusion proteins are the

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Mal-E fusion proteins and QIAexpress system fusion proteins (QIAGEN, USA) as detailed above. The fusion or mixed protein may include, either instead of in addition to the urease sub-unit, a Heat Shock Protein, or fragment or variant thereof, as defined above.

The invention also relates to monoclonal polyclonal antibodies to the proteinaceous materials described above. More particularly, the invention relates to antibodies or fragments thereof to any one of the Helicobacter felis polypeptides encoded by the urease gene cluster of the plasmid pILL205 (CNCM I-1355) including the structural and accessory urease polypeptides that is, structural genes ure A and ure B and the accessory genes known as ure C, ure D, ure E, ure F, ure G, ure H and ure I. The antibodies may also be directed to a polypeptide having at least 90 % homology with any of the above urease polypeptides or to a fragment thereof preferably having at least 6 The antibodies of the invention may amino-acids. specifically recognise Helicobacter felis polypeptides expressed by the urease gene cluster. In this case, the epitopes recognised by the antibodies are unique to Helicobacter felis. Alternatively, the antibodies include or consist of antibodies directed to epitopes common Helicobacter felis to polypeptides and to Helicobacter pylori Ιf polypeptides. the antibodies recognise the accessory gene products, it is particularly advantageous that they cross-react with the Helicobacter pylori accessory gene product. In this antibodies may be used in therapeutic treatment of Helicobacter pylori infection in man, by blocking the urease maturation process.

Particularly preferred antibodies of the invention recognise the <u>Helicobacter felis</u> ure A

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and/or <u>ure B</u> gene products, that is the A and B urease sub-units. Advantageously, these antibodies also cross-react with the <u>Helicobacter pylori</u> A and B urease sub-units, but do not cross-react with other ureolytic bacteria. Such antibodies may be prepared against epitopes unique to <u>Helicobacter</u> (see figure 4), or alternatively, against the whole polypeptides followed by screening out of any antibodies reacting with other ureolytic bacteria.

The invention also concerns monoclonal or polyclonal antibodies to the HSPs or fragments thereof, particularly to the HSP A and/or HSP B protein illustrated in figure 6. Polypeptides having at least 75 %, and preferably at least 80 %, or 90 % homology with the HSPs may also be used to induce antibody formation. These antibodies may be specific Helicobacter pylori chaperonins alternatively, they may cross-react with GroEL-like proteins or GroES-like proteins from bacteria other than <u>Helicobacter</u>, depending upon the epitopes recognised. Figure 7 shows the homologous regions of HSP A and HSP B with GroES-like proteins and GroELlike proteins respectively from various bacteria. Particularly preferred antibodies are those specific for either the HSP A or HSP B chaperonins or those specifically recognising the HSP A C-terminal sequence having the metal binding function. Again, specific fragments for the induction of the antibodies ensures production of Helicobacter-specific antibodies.

The antibodies of the invention may be prepared using classical techniques. For example monoclonal antibodies may be produced by the hybridoma technique or by known techniques for the preparation of human antibodies, or by the technique described by Marks et

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al (Journal of Molecular Biology, 1991, 222, p 581-597).

The invention also includes fragments of any of the above antibodies produced by enzyme digestion. Of particular interest are the Fab and F(ab')₂ fragments. Also of interest are the Facb fragments.

The invention also relates to purified antibodies or serum obtained by immunisation of an animal, e.g. a the immunogenic composition, mammal, with proteinaceous material or fragment, or the fusion or of the invention, protein followed by purification of the antibodies or serum. concerned is a reagent for the in vitro detection of H. pylori infection, containing at least antibodies or serum, optionally with reagents for labelling the antibodies e.g. anti-antibodies etc.

The invention further relates to nucleic acid sequences coding for any of the above proteinaceous materials including peptides. In particular, the invention relates to a nucleic acid sequence characterised in that it comprises:

- i) a sequence coding for the <u>Helicobacter felis</u> urease and accessory polypeptides as defined above, and a sequence coding for the HSP of <u>H. pylori</u> as defined above;
- or ii) a sequence complementary to sequence (i); or iii) a sequence capable of hybridizing to sequence (i) or (ii) under stringent conditions; or iv) a fragment of any of sequences (i), (ii) or (iii) comprising at least 10 nucleotides.

Preferred nucleic acid sequences are those comprising all or part of the sequence of plasmid pIL205 (CNCM I-1355), for example the sequence of Figure 3, in particular that coding for the gene product of <u>ure A</u> and for <u>ure B</u> or the sequence of

Figure 9 (<u>Ure I</u>), or a sequence capable of hybridising with these sequences under stringent conditions, or a sequence complementary to these sequences, or a fragment comprising at least 10 consecutive nucleotides of these sequences.

Other preferred sequences are those comprising all or part of the sequence of plasmid pILL689 (CNCM I-1356), for example the sequence of figure 6, in particular that coding for HSP A and/or HSP B, or a sequence complementary to this sequence, or a sequence capable of hybridizing to this sequence under stringent conditions, or a fragment thereof.

High stringency hybridization conditions in the context of the invention are the following:

- 5 x SSC ;
- 50 % formamide at 37°C; or:
 - 6 x SSC ;
 - Denhard medium at 68°C.

The sequences of the invention also include those hybridizing to any of sequences (i), (ii) and (iii) defined above under non-stringent conditions, that is:

- 5 x SSC ;
- 0.1 % SDS ;
- 30 or 40 % formamide at 42°C, preferably 30 %.

The term "complementary sequences" in the context of the invention signifies "complementary" and "reverse" or "inverse" sequences.

The nucleic acid sequences may be DNA or RNA.

The sequences of the invention may be used as nucleotide probes in association with appropriate labelling means. Such means include radio-active isotopes, enzymes, chemical or chemico-luminescent markers, fluoro-chromes, haptens, or antibodies. The

markers may optionally be fixed to a solid support, for example a membrane, or particles.

As a preferred marker, radio-active phosporous (32P) is incorporated at the 5'-end of the probe sequence. The probes of the invention comprise any fragment of the described nucleic acid sequences and may have a length for example of at least 45 nucleotides, for example 60, 80 or 100 nucleotides or more. Preferred probes are those derived from the ure A, ure B, ure I, HSP A and HSP B genes.

The probes of the invention may be used in the in vitro detection of Helicobacter infection optionally biological sample, after gene amplification reaction. Most advantageously, the probes are used to detect Helicobacter felis Helicobacter pylori, or both, depending on whether the sequence chosen as the probe is specific to one or the other, or whether it can hybridise to both. Generally, the hybridisation conditions are stringent in carrying out such a detection.

The invention also relates to a kit for the <u>in</u>

<u>vitro</u> detection of <u>Helicobacter</u> infection,

characterised in that it comprises:

- a nucleotide probe according to the invention, as defined above;
- an appropriate medium for carrying out a hybridisation reaction between the nucleic acid of <u>Helicobacter</u> and the probe;
- reagents for the detection of any hybrids formed.

The nucleotide sequences of the invention may also serve as primers in a nucleic acid amplification reaction. The primers normally comprise at least 10 consecutive nucleotides of the sequences described above and preferably at least 18. Typical lengths are

from 25 to 30 and may be as high as 100 or more consecutive nucleotides. Such primers are used in pairs and are chosen to hybridize with the 5' and 3'fragment to be amplified. ends of the amplification reaction may be performed using for technique example the PCR (European applications EP200363, 201184 and 229701). The $Q-\beta$ replicase technique (Biotechnology, vol. 6, Oct. 1988) may also be used in the amplification reaction.

The invention also relates to expression vectors characterised in that they contain any of the nucleic acid sequences of the invention. Particularly preferred expression vectors are plasmids pILL689 and pILL205 (CNCM I-1356 and CNCM I-1355, respectively). The expression vectors will normally contain suitable promoters, terminators and marker genes, and any other regulatory signals necessary for efficient expression.

The invention further relates to prokaryotic or eukaryotic host cells stably transformed by nucleic acid sequences of the invention. As examples of hosts, mention may be made of higher eukaryotes such as CHO cells and cell-lines; yeast, prokaryotes including bacteria such as E. coli e.g E. coli HB 101 Mycobacterium tuberculosum ; viruses baculovirus and vaccinia. Usually the host cells will be transformed by vectors. However, it possible within the context of the invention, insert the nucleic acid sequences by homologous recombination, using conventional techniques.

By culturing the stably transformed hosts of the invention, the <u>Helicobacter</u> urease polypeptide material and, where applicable, the HSP material can be produced by recombinant means. The recombinant proteinaceous materials are then collected and purified. Pharmaceutical compositions are prepared by

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combining the recombinant materials with suitable excipients, adjuvants and optionally, any other additives such as stabilizers.

The invention also relates to plasmids pILL920 (deposited at CNCM on 20.07.1993, under accession number I-1337) and pILL927 (CNCM I-1340, deposited on 20.07.1993) constructed as described in the examples below.

Different aspects of the invention are illustrated in the figures:

Figure 1:

Transposon mutagenesis and sequencing of pILL205. Linear restriction maps of recombinant cosmid pILL199 and recombinant plasmid pILL205 (and the respective scale markers) are presented. Numbers in parentheses indicate the sizes of H.felis DNA fragments inserted into one of the cloning vectors (pILL575 or pILL570, respectively). The "plus" and "minus" signs within circles correspond to the insertion sites of MiniTn3-Km transposon in pILL205 ; "plus" indicate that the transposon did not inactivate urease expression, whereas negative signs indicate that urease expression was abolished. The letters refer to mutant clones which were further characterised for quantitative urease activity and for the synthesis of urease gene products. The location of the structural urease genes (ure A and ure B) on pILL205 represented by boxes, the lengths of which proportional to the sizes of the respective openreading frames. The arrows refer to the orientation of transcription. The scale at the bottom of the figure indicates the sizes (in kilobases) of the HindIII and PstI restriction fragments. Restriction sites

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represented as follows: B, BamHI; Pv, PvuII; Bg, BglII; E, EcoRI; H, HindIII; C, ClaI; Ps, PstI. Letters within parentheses indicate that the sites originated from the cloning vector.

Figure 2:

Western blot analysis of whole-cell extracts of E. coli HB101 cells harbouring recombinant plasmids were reacted with rabbit polyclonal antiserum (diluted 1000) raised against H. felis bacteria. extracts were of E. coli cells harbouring : plasmid vector pILL570 (lane 1); recombinant plasmid pILL205 (lane 2); and pILL205 derivative plasmids disrupted in loci "a", "b", "c", "d", and "e" (lanes 3-7). B) Extracts were of E. coli cells harbouring : recombinant plasmid pILL753 containing the H. pylori ure A and ure B genes (Labigne et al., 1991) (lane 1) ; and pILL205 derivative plasmids disrupted in loci "f", "g", "h", and "i" (lanes 2-5). The small arrow heads indicate polypeptides of approximately 30 and 66 kilodaltons which represent putative Ure A and Ure B gene products of H. felis. The large arrow heads in panel B indicate the corresponding gene products of H. pylori which cros-reacted with the anti-H. felis serum. The numbers indicate the molecular weights (in thousands) of the protein standards.

Figure 3:

Nucleotide sequence of the <u>H. felis</u> structural urease genes. Numbers above the sequence indicate the nucleotide positions as well as the amino acid position in each of the two <u>Ure A</u> and <u>Ure B</u> polypeptides. Predicted amino acid sequences for <u>Ure A</u> (bp 43 to 753) and <u>Ure B</u> (766 to 2616) are shown below

the sequence. The putative ribosome-binding site (Shine-Dalgarno sequence, SD) is underlined.

Figure 4:

Comparison of sequences for the structural urease genes of H. felis to : a) the sequence of the two subunits of H. pylori urease (Labigne et al., 1991); b) the sequence of the three subunits of Proteus mirabilis urease (Jones and Mobley, 1989); c) the sequence of the single subunit of jack bean urease. Gaps (shown by dashes) have been introduced to ensure the best alignment. *, amino acids identical to those of the H. felis sequence ; =, amino-acids shared by the various ureases; , amino-acids unique to the Helicobacter ureases. The percentages relate to the number of amino acids that are identical to those of the <u>H. felis</u> urease subunits. <u>H.f.</u>, Helicobacter H.p., Helicobacter pylori; P.m., Proteus felis ; mirabilis ; J.b., Jack bean.

Figure 5 :

Restriction map of the recombinant plasmids pILL689, pILL685, and pILL691. The construction of these plasmids is described in details in Table 1. Km within triangles depictes the site of insertion of the kanamycin cassette which led to the construction of plasmids pILL687, pILL688 and pILL696 (table 2). Boxes underneath the maps indicate the position of the three genetic elements deduced from the nucleotide sequence, namely IS5, Hsp A and Hsp B.

Figure 6:

Nucleotide sequence of the <u>Helicobacter pylori</u>
Heat Shock Protein gene cluster. The first number above the sequence indicates the nucleotide positions, whereas the second one numbers the amino-acid residue

position for each of the <u>Hsp A</u> and <u>Hsp B</u> protein. The putative ribosome-binding sequences (Shine- Dalgarno [SD] sites) are underlined.

Figure 7:

Comparison of the deduced amino-acid sequence of <u>Helicobacter pylori</u> <u>Hsp A</u> (A) or <u>Hsp B</u> (B) with that of other GroEL-like (A) or GroES-like (B) proteins. Asteriks mark amino-acids identical with those in the <u>Helicobacter pylori Hsp A</u> or <u>Hsp B</u> sequences.

Figure 8:

Expression of the <u>Helicobacter pylori Hsp A</u>
Heat-Shock proteins in <u>E. coli</u> minicells. The protein bands with apparent molecular masses of 58 and 13 kDA, corresponding to the <u>Helicobacter pylori Hsp A</u> and <u>Hsp B</u> Heat-Shock Proteins are clearly visible in the lanes corresponding to plasmids pILL689 and pILL692 and absent in the vector controls (pILL570 and pACYC177, respectively)

Figure 9 :

Nucleotide sequence of the <u>Helicobacter felis</u> <u>ure</u> <u>I</u> gene and deduced amino-acid sequence.

Figure 10 :

Comparison of the amino-acid sequence of the <u>ure</u> I proteins deduced from the nucleotide sequence of the <u>ure I gene of Helicobacter felis</u> and that of Helicobacter pylori.

Figure 11:

Genetic code. Chain-terminating, or "nonsense", codons. Also used to specify the initiator formyl-Met-tRNA $^{Met}_{\rm F}$. The Val triplet GUG is therefore

"ambiguous" in that it codes both valine and methionine.

Figure 12:

Signification of the one-letter and three-letter amino-acid abbreviations.

Figure 13 :

Purification of H. pylori UreA-MBP recombinant protein using the pMAL expression vector Extracts from the various stages of protein purification were migrated on a 10 % resolvving SDSpolyacrylamide gel. Following electrophoresis, the gel was stained with Coomassie blue. The extracts were: 1) non-induced cells; 2) IPTG-induced cells; French press lysate of induced cell extract; 5) eluate from amylose resin column; 6) eluate from anion exchange column (first passage) ; 7) eluate from anion exchange column (second passage); 8) SDS-PAGE standard marker proteins.

Figure 14:

Recognition of UreA recombinant fusion proteins by polyclonal rabbit anti-Helicobacter sera. Protein extracts of maltose-binding protein (MBP, lane 1), H. felis UreA-MBP (lane 2), and H. pylori UreA-MBP (lane 3) were Western Blotted using rabbit polyclonal antisera (diluted 1:5000) raised against whole-cell extracts of H. pylori and H. felis. The purified fusion proteins are indicated by an arrow. Putative degradation products of the proteins are shown by an asterisk.

Figur 15:

Recognition of UreB recombinant fusion proteins by rabbit antisera raised against purified homologous

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and heterologous UreB proteins. Nitrocellulose membranes were blotted with the following extracts:

1) standard protein markers; 2) H. felis UreA-MBP;

3) MBP; 4) H. pylori UreA-MBP. The membranes were reacted with polyclonal rabbit antisera (diluted 1:5000) raised against MBP-fused H. pylori and H. felis Ure B sub-units, respectively. The molecular weights of standard proteins are presented on the left-hand side of the blots.

Figure 16:

Western blot analysis of <u>H. pylori</u> and <u>H. felis</u> whole-cell extracts with antisera raised against purified UreB MBP-fused recombinant proteins. SDS-PAGE whole extracts of <u>H. Felis</u> (lane 1) and <u>H. pylori</u> (lane 2) cells were reacted with polyclonal rabbit antisera raised against purified <u>H. pylori</u> UreB and <u>H. felis</u> UreB MBP-fused proteins (sera diluted 1 : 5000). The difference in gel mobility of the respective non-recombinant UreB sub-units of <u>H. felis</u> and <u>H. pylori</u> can be seen. The numbers on the left refer to the molecular weights of standard marker proteins.

Figure 17:

SDS-PAGE analysis of material eluted from the amylose column (lanes 2 and 3) or from the Ni-NTA column following elution: *with buffer E (pH 4.5), lanes 4 and 5; or buffer C (pH 6.3), lanes 6 and 7. Material eluted from a lysate of MC1061 (PILL933) (lanes 2, 3, 5 and 7) and material eluted from a lysate of MC1061 (PMAL-c2) (lanes 4 and 6). Lane 3 contains the same material as in lane 2 except that it was resuspended in buffer E, thus demonstrating that buffer E is responsible for dimer formation of the MBP-HspA subunit, as seen in lanes 3 and 5.

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Figure 18 :

Serum IgG responses to MBP (bottom), MBP-HspA (top) and MBP-HspB (middle) of 28 <u>H. pylori</u> infected patients (squares, left) and 12 uninfected patients (circles, right). The optical density of each serum in the ELISA assay described in Experimental procedures was read at 492 nm, after a 30 mn incubation. The sizes of the symbols are proportional to the number of sera giving the same optical density value.

EXAMPLES

I - CLONING, EXPRESSION AND SEQUENCING OF H. FELIS UREASE GENE :

EXPERIMENTAL PROCEDURES FOR PART I:

Bacterial strains and culture conditions :

H. felis (ATCC 49179) was grown on blood agar base no. 2 (Oxoid) supplemented with 5 % (v/v) lysed horse blood (BioMerieux) and an antibiotic supplement ml' vancomycin (Lederle consisting of 10 ng Laboratories), 2.5 μg ml⁻¹ polymyxin B (Pfizer), 5μg ml-1 trimethoprim (Sigma Chemical Co.) and 2.5 μg ml-1 amphotericin B (E.R Squibb and Sons, Inc.). Bacteria were cultured on freshly prepared agar plates and lid uppermost, incubated, under microaerobic conditions at 37°C for 2-3 days. E. coli strains HB101 Roulland-Dussoix, 1969) (Boyer and and MC1061 al., 1983), (Maniatis et used in the experiments, were grown routinely in Luria broth without glucose added or on Luria agar medium, 37°C. Bacteria under grown nitrogen-limiting conditions were passaged on a nitrogen-limiting solid medium consisting of ammonium-free M9 minimal medium (pH 7.4) supplemented with 0.4 % (w/v) D-glucose and 10 mM L-arginine (Cussac et al., 1992).

DNA manipulations :

All standard DNA manipulations and analyses, unless mentioned otherwise, were performed according to the procedures described by Maniatis et al. (1983).

Isolation of H. felis DNA:

Total genomic DNA was extracted by an sarkosylproteinase K lysis procedure (Labigne-Roussel et al., 1988). Twelve blood agar plates inoculated with \underline{H} . felis were incubated in an anaerobic jar (BBL) with an anaerobic gaspak (BBL 70304) without catalyst, for 1-2 days at 37°C. The plates were harvested in 50 ml of a 15 % (v/v) glycerol - 9 % (w/v) sucrose solution and centrifuged at 5,000 rpm (in a Sorvall centrifuge), for 30 min at 4°C. The pellet was resuspended in 0.2 ml 50 mM D-glucose in 25 mM Tris-10 mM EDTA (pH 8.0) containing 5 mg ml⁻¹ lysozyme and transferred to a VTi65 polyallomer quick seal tube. A 0.2 ml aliquot of 20 mg ml^{-1} proteinase K and 0.02 ml of 5M sodium perchlorate were added to the suspension. Cells were lysed by adding 0.65 ml of 0.5M EDTA -10 % (W/V)Sarkosyl, and incubated at 65°C until the suspension cleared (approximately 5 min). The volume of the tube was completed with a CsCl solution consisting (per 100 ml) of 126 g CsCl, 1 ml aprotinine, 99 ml TES buffer (30 mM Tris, 5 mM EDTA, 50 mM NaCl (pH 7.5). Lysates were centrifuged at 45 000 rpm, for 15-18 h at 18°C. was collected and dialysed against TE Total DNA buffer (10 mM Tris, 1 mM EDTA), at 4°C.

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Cosmid cloning:

Chromosomal DNA from H. felis was cloned into vector pILL575, as previoulsy described (Labigne et al, 1991). Briefly, DNA fragments arising from a partial digestion with Sau3A were sized on a (10 to 40 %) sucrose density gradient and then ligated into a BamHI-digested and dephosphorylated pILL575 DNA preparation. Cosmids were packaged into phage lambda particles (Amersham, In Vitro packaging kit) and used infect E. coli HB101. To screen for urease expression, kanamycin-resistant transductants replica-plated onto solid nitrogen-mimiting medium (see above) containing (20 μ g ml⁻¹) kanamycin that had been dispensed into individual wells of microtitre plates (Becton Dickinson). The mictrotitre plates were incubated aerobically, at 37°C for 2 days before adding 0.1 ml urease reagent (Hazell et al., 1987) to each of the wells. Ureolysis was detected within 5-6 h at 37°C by a colour change in the reagent. Several urease-positive cosmid clones were restriction mapped and one was selected for subcloning.

Subcloning of H. felis DNA:

A large-scale CsCl plasmid preparation of cosmid DNA was partially digested Sau3A. DNA fragments (7 - 11 kb) were electroeluted from an agarose gel and purified using phenol-chloroform extractions. Following precipitation in cold ethanol, the fragments were ligated into Bg/III-digested plasmid pILL570 (Labigne et al., 1991) and the recombinant plasmids used to transform competent E. coli MC1061 cells. Spectinomycin-resistant transformants were selected and screened for urease expression under nitrogen-rich (Luria agar) and nitrogen-limiting conditions.

Quantitative urease activity:

Cultures grown aerobically for 2.5 days at 37°C were harvested and washed twice in 0.85 % (w/v) NaCl. Pellets were resuspended in PEB buffer (0.1 M sodium phosphate buffer (pH 7.4) containing 0.01 M EDTA) and then sonicated by four 30-sec bursts using a Branson Sonifier model 450 set at 30 W, 50 % cycle. Cell debris was removed from the sonicates centrifugation. Urease activities of the sonicates were measured in a 0.05 M urea solution prepared in PEB by a modification of the Berthelot reaction (Cussac et al., 1992). Urease activity was expressed as μ mol urea min⁻¹mg⁻¹ bacterial protein.

Protein determination :

Protein concentrations were estimated with a commercial version of the bradford assay (Sigma Chemicals).

Transposon mutagenesis:

Random insertional mutations were generated within cloned <u>H. felis</u> via a MiniTn3-Km delivery system (Labigne et al., 1992). In brief, <u>E. coli</u> HB101 cells containing the transposase-encoding plasmid pTCA were transformed with plasmid pILL570 containing cloned <u>H. felis</u> DNA. Transposition of the MiniTn3-Km element into the pILL570 derivative plasmids was effected via conjugation. The resulting cointegrates were then selected for resolved structures in the presence of high concentrations of kanamycin (500 mg1-1) and spectinomycin (300 mg1-1).

SDS-PACE and Western blotting :

Solubilised cell extracts were analysed on slab gels, comprising a 4.5 % acrylamide stacking gel and 12.5 % resolving gel, according to the procedure of

Laemmli (Laemmli, 1970). Electrophoresis was performed at 200V on a mini-slab gel apparatus (Bio-Rad).

Proteins were transferred to nitrocellulose paper (Towbin et al., 1979) in a Mini Trans-Blot transfer cell (Bio-Rad) set at 100 V for 1 h (with cooling). Nitrocellulose membranes were blocked with 5 % (w/v)purified casein (BDH) in phosphate-buffered saline (PBS, pH 7.4) at room temperature, for 2 h (Ferrero et al., 1992). Membranes were reacted at 4°C overnight with antisera diluted in 1 % (w/v) casein prepared in Immunoreactants were then detected biotinylated secondary antibody (Kirkegaard and Perry Lab.) in combination with avidin-peroxidase (KPL). A substrate solution composed of 0.3 % (w/v)chloro-1-naphthol (Bio-rad) was used to visualise reaction products.

DNA Sequencing:

DNA fragments to be sequenced were cloned into M13mp18 and M13mp19 (Meissing and Vieira, 1982) bacteriophage vectors (Pharmacia). Competent <u>E. coli</u> JM101 cells were transfected with recombinant phage DNA and plated on media containing X-gal (5-bromo-4-chloro-3-indolyl- β -D-galactopyranoside) and isopropyl- β -D-thiogalactopyranoside. Plaques arising from bacteria infected with recombinant phage DNA were selected for the preparation of single-stranded DNA templates by polyethylene glycol treatment (Sanger et al., 1977). Single-stranted DNA sequenced according to the dideoxynucleotide chain termination method using a Sequenase kit (United States Biochemical Corp.).

Nucleotide sequence acc ssion number :

The nucleotide accession number is X69080 (EMBL Data Library).

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RESULTS OF PART I EXPERIMENTS :

Expression of urease activity by H. felis cosmid clones:

Cloning of partially digested fragments (30 to 45 kb in size) of H. felis chromosomal DNA into the cosmid vector pILL575 resulted in the isolation of approximately 700 cosmid clones. The clones were subcultured on nitrogen-limiting medium in order to induce urease expression (Cussac et al., 1992). Six of these were identified as being urease-positive after 5-6 h incubation (as described in the Experimental procedures section). No other urease-positive cosmid clones were identified, even after a further overnight incubation. Restriction enzyme analysis of 3 clones harbouring the urease-encoding cosmids revealed a 28 kd DNA fragment. A cosmid (designated pILL199) containing DNA regions at both extremities of the common fragment was selected for subcloning.

Identification of H. felis genes required for urease expression when cloned in E. coli cells:

To define the minimum DNA region necessary for urease expression in <u>E. coli</u> cells, the urease-encoding cosmid pILL199 was partially digested with Sau3A and the fragments were subcloned into plasmid pILL570. The transformants were subcultured on nitrogen-rich and nitrogen-limiting media and screened for an urease-positive phenotype. Five transformants expressed urease activity when grown under nitrogen-limiting conditions, whereas no activity was detected following growth on nitrogen-rich medium. Restriction mapping analyses indicated that the urease-encoding plasmids contained inserts of between 7 and 11 kb. The

plasmid designated pILL205 was chosen for further studies.

Random mutagenesis of cloned H. felis DNA was performed to investigate putative regions essential for urease expression in E. coli and to localise the region of cloned DNA that contained the structural Random insertion mutants urease genes. prototype plasmid pILL205 were thus generated using the MiniTn3-Km element (Labigne et al, 1992). The site of insertion was restriction mapped for each of the mutated copies of pILL205 and cells harbouring these plasmids were assessed qualitatively for activity (figure 1). A selection of E. coli HB101 cells harbouring the mutated derivatives of pILL205 (designated "a" to "i") were then used both for quantitative urease activity determinations, as well as for the detection of the putative urease subunits by Western blotting.

The urease activity of E. coli HB101 cells harbouring pILL205 was 1.2 \pm 0.5 μ mol urea min⁻¹mg⁻¹ bacterial protein (table 1), which is approximately a fifth that of the parent H. felis strain used for the cloning. Insertion of the transposon at sites "a", "d", "f" and "g" resulted in a negative phenotype, whilst mutations at sites "b", "e", "h" and "i" had no significant effect on the urease activities of clones harbouring these mutated copies of pILL205 (table 1). Thus mutagenesis of pILL205 with the MiniTn3-Km element identified three domains as being required for \underline{H} . felis wrease gene expression in \underline{E} . coli cells.

Localisation of the H. f lis urease structural genes :

Western blot analysis of extracts of \underline{E} . \underline{coli} cells harbouring pILL205 indicated the presence of two

polypeptides of approximately 30 and 66 kDa which cross-reacted with polyclonal H. felis antiserum (Figure 2A). These proteins were produced by bacteria carrying the vector (pILL570). Native H. felis urease has been reported to be composed of repeating monomeric subunits calculated molecular weights of 30 and 69 kDa (Turbett et al, 1992). Thus the 30 and 66 kDa proteins were thought to correspond to the ure A and ure B gene products, respectively. Interestingly an extract of E. coli cells harbouring the recombinant plasmid pILL763 1992) containing the <u>Helicobacter</u> (Cussac et al, pylori ure A and ure B genes, expressed polypeptides with approximate molecular sizes of 30 and 62 kDa which cross-reacted with the anti-H. felis antisera (figure 2B).

Table 1. Mutagenesis of E. coli clones and effect on urease activity.

plasmids ^a	Urease activity ^b
	(µmol urea min-1 mg-1 protein)
pILL205	1.2 ± 0.46 °
pILL205 :: a	neg ^d
pILL205":: b	0.74 ± 0.32
pILL205 :: c	neg
pILL205 :: d	neg
pILL205 :: e	0.54 ± 0.15
pILL205 :: f	neg
pILL205 :: g	neg
pILL205 :: h	1.05 ± 0.25
pILL205 :: i	0.93 ± 0.35

- E. coli cells harboured pILL205 and its derivatives constructed by transposon mutagenesis. The letters correspond to the insertion sites of the MiniTn3-transposon on pILL205.
- Activities of bacteria grown aerobically for 3 days at 37 °C on solid M9 minimal medium supplemented with 10 mM L-arginine. The values represent the means ± standard deviations calculated from three determinations.
- Urease activity was approximately a fifth as large as that of H. felis wild-type strain (ATCC 49179) i.e. $5.7 \pm 0.1 \, \mu mol \ urea \ min^{-1} \ mg^{-1}$ protein (Ferrero and Lee, 1991).
- d No activity detected (limit of detection was < 1 nmol urea min⁻¹ mg⁻¹ of bacterial protein).

Clones harbouring the mutated derivatives of pILL205, in all but one case, expressed the ure A and ure B gene products (Figures 2A, B). Given that several of the mutants (i.e. mutants "c", "d", "f" and "g") synthesised the urease subunits yet did not produce an active enzyme, it is possible to speculate that accessory functions essential for urease activity may have been disrupted by transposon insertion. In contrast, the mutant designated pILL205::a did not produce the <u>ure B</u> product and was urease-negative. Thus the site of transposon insertion was presumed to be located in the ure B gene. Sequence analyses of the DNA region corresponding to insertion site "a" were undertaken to elucidate potential open reading frames encoding the structural polypeptides of H. felis urease.

<u>Bequence analyses of H. felis structural urease</u> genes:

sequencing of a 2.4 kb region of H. felis DNA adjacent to transposon insertion site "a" resulted in the identification of two open reading frames (ORFs) designated <u>ure A</u> and <u>ure B</u> which are transcribed in the same direction (figure 3). The transposon was confirmed to be located at 240 bp upstream from the end of <u>ure B</u>. Both ORFs commenced with an ATG start codon and were preceded by a site similar to the <u>E</u>. coli consensus ribozome-binding sequence (Shine and Dalgarno, 1974). The intergenic space for the <u>H</u>. felis structural genes consisted of three codons which were in phase with the adjacent open-reading frames. This suggests that, as has already been observed to be the cas for <u>Helicobacter pylori</u> (Labique et al, 1991), a single mutation in the stop codon of the <u>ure A</u> gene

would theoretically result in a fused single polypeptide.

The H. felis ure A and ure B genes encode polypeptides with calculated molecular weights of 26 074 kA and 61 663 Da, respectively, which are highly homologous at the amino-acid sequence level to the ure A and ure B gene products of H. pylori. The levels of identity between the corresponding ure A and ure B gene products of the two Helicobacter spp. was calculated to be 73.5 % and 88.2 % respectively. From the amino-acid sequence information, the predicted molecular weights of the ure A and ure B polypeptides from H. felis and H. pylori (Labigne et al, 1991) are very similar. Nevertheless the ure B product of H. felis had a lower mobility than the corresponding gene product from Helicobacter pylori when subjected to SDS-polyacrylamide gel electrophoresis (figure 2B)

II - EXPRESSION OF RECOMBINANT UREASE SUBUNIT PROTEINS FROM H. PYLORI AND H. FELIS: ASSESSMENT OF THESE PROTEINS AS POTENTIAL MUCOSAL IMMUNOGENS IN A MOUSE MODEL:

The aims of the study were to develop recombinant antigens derived from the urease subunits of <u>H. pylori</u> and <u>H. felis</u>, and to assess the immunoprotective efficacies of these antigens in the <u>H. felis</u>/mouse model. Each of the structural genes encoding the respective urease subunits from <u>H. pylori</u> and <u>H. felis</u> was independently cloned and over-expressed in <u>Escherichia coli</u>. The resulting recombinant urease antigens (which were fused to a 42 kDa maltose-binding protein of <u>E. coli</u>) were purified in large quantities from <u>E. coli</u> cultures and were immunogenic, yet enzymatically inactive. The findings demonstrated the

feasibility of developing a recombinant vaccine against <u>H. pylori</u> infection.

EXPERIMENTAL PROCEDURES FOR PART II :

Bacterial strains, plasmids and growth conditions:

H. felis (ATCC 49179) was grown on a blood agar medium containing blood agar base no. 2 (Oxoid) supplemented with 10% lysed horse blood (BioMérieux) and an antibiotic supplement consisting of vancomycin (10 μ g/mL), polymyxin B (25 ng/mL), trimethoprim (5 μ g/mL) and amphotericin B (2.5 μ g/mL). Bacteria were cultured under microaerobic conditions at 37° C for 2 days, as described previously. E. coli strains MC1061 and JM101, used in cloning and expression experiments, were grown routinely at 37° C in Luria medium, with or without agar added. The antibiotics carbenicillin (100 μ g/mL) and spectinomycin (100 μ g/mL) were added as required.

DNA manipulations and analysis:

manipulations and analyses, All DNA otherwise, were performed according mentioned procedures. Restriction and modification standard enzymes were purchased from Amersham (France). DNA fragments to be cloned were electroeluted from agarose gels and then purified by passage on Elutip mini-(Schleicher and Schull, Germany). columns stranded DNA sequencing was performed using M13mpl8 and M13mp19 bacteriophage vectors (Pharmacia, France). Single-stranded DNA templates were prepared phage DNA by polyethylene recombinant glycol treatment. Sequencing of the templates was achieved according to the dideoxynucleotide chain termination method using a Sequenase kit (United States Biochemical Corp., U.S.A.).

Preparation of inserts for cloning using the polymerase chain reaction (PCR):

To clone the ureA genes of H. pylori and H. felis, degenerated 36-mer primers were conceived from the published urease sequences (Labigne et al., 1991; Ferrero and Labigne, 1993) (primer set #1 ; refer to table 2). Purified DNA from E. coli clones harbouring plasmids pILL763 and pILL207 (table 3), that encoded the structural genes of H. pylori and H. felis ureases, were used as template material in reactions. Reaction samples contained: 10 - 50 ng of denatured DNA; PCR buffer (50 mmol/L KCl in 10 mmol/L Tris-HCl [pH 8.3)]); dATP, dGTP, dCTP and dTTP (each at a final concentration of 1.25 mmol/L) ; 2.5 mmol/L MgCl,; 25 pmol of each primer and 0.5 µL Tag polymerase. The samples were subjected to 30 cycles of the following programme : 2 min at 94° C, 1 min at 40° c.

The amplification products were cloned into the cohesive ends of the pAMP vector (figure 1) according protocol described by the manufacturer System", Gibco BRL ; Cergy ("CloneAmp France). Briefly, 60 ng of amplification product was directly mixed in a buffer (consisting of 50 mmol/L KCl, 1.5 mmol/L MgCl₂, 0.1 % (wt/vol) gelatine in 10 mmol/L Tris-HCl, pH 8.3) with 50 ng of the pAMP 1 vector DNA and 1 unit of uracil DNA glycolsylase. Ligation was performed for 30 min at 37° C. Competent cells (200 μ L) of E. coli MC1061 were transformed with ligation mixture. Inserts 20 μ L the subsequently excised from the polylinker of the pAMP vector by double digestion with BamH1 and Pst1, and then subcloned into the expression vector pMAL (New England Biolabs Inc., Beverly, USA) chosen for the production of recombinant antigens (pILL919 and pILL920, respectively, figure 13), as well as in M13mp bacteriophage for sequencing.

Amplification of a product containing the <u>ureB</u> gene of <u>H. pylori</u> was obtained by PCR using a couple of 35-mer primers (set #2, table 2). The PCR reaction mixtures were first denatured for 3 min at 94° C, then subjected to 30 cycles of the following programme: 1 min at 94° C, 1 min at 55° C and 2 min at 72° C. The purified amplification product (1850 bp was digested with <u>EcoRI</u> and <u>PstI</u> and then cloned into pMAL (pILL927, figure 2). Competent cells of <u>E. coli</u> MC1061 were transformed with the ligation reaction.

H. felis ureB was cloned in a two-step procedure, that allowed the production of both complete and truncated versions of the UreB subunit. pILL213 (table 3) was digested with the enzymes DraI, corresponding to amino acid residue number 219 of the The resulting 1350 UreB subunit and HindIII. fragment was purified and cloned into pMAL that had been digested with XmnI and HindIII (pILL219, figure order to produce a capable clone 2). synthesizing a complete UreB protein, PCR primers were developed (set #3, table 2), that amplified a 685 bp fragment from the N-terminal portion of the ureB gene (excluding the ATG codon), that also overlapped the beginning of the insert in plasmid pILL219. The PCR amplified material was purified and digested with bamHI and HindIII, and then cloned into pMAL (pILL221, figure 14). A 1350 bp PstI-PstI fragment encoding the remaining portion of the UreB gene product was subsequently excised from pILL219 and cloned into a linearised preparation of pILL221 (pILL222, figure 14).

Expression of recombinant urease polypeptides in the vector pMAL:

The expression vector pMAL is under the control of an inducible promoter (P_{lac}) and contains an open-reading frame (ORF) that encodes the production of MalE (Maltose-binding protein, MBP). Sequences cloned in-phase with the latter ORF resulted in the synthesis of MBP-fused proteins which were easily purified on amylose resin. Of the two versions of pMAL that are commercially available, the version not encoding a signal sequence (ie. pMAL-c2) synthesized greater amounts of recombinant proteins and was thus used throughout.

<u>E. coli</u> clones harbouring recombinant plasmids were screened for the production of fusion proteins, prior to performing large-scale purification experiments.

Purification of recombinant urease polypeptides:

Fresh 500 mL volumes of Luria broth, containing carbenicillin (100 μ g/mL and 2% (wt/vol) glucose, were inoculated with overnight cultures (5 mL) of <u>E. colictorial</u> clones. The cultures were incubated at 37° C and shaken at 250 rpm, until the $A_{600} = 0.5$. Prior to adding 1 mmol/L (final concentration) isopropyl- β -D-thiogalactopyranoside (IPTG) to cultures, a 1.0 mL sample was taken (non-induced cells). Cultures were incubated for a further 4 h at which time another 1.0 mL sample (induced cells) was taken. The non-induced and induced cell samples were later analysed by SDS-PAGE.

IPTG-induced cultures were centrifuged at 7000 rpm for 20 min, at 4° C and the supernatant discarded. Pellets were resuspended in 50 mL column buffer (200 mmol/L NaCl, 1 mmol/L EDTA in 10 mmol/L TrisHCl,pH 7.4), containing the following protease inhibitors (supplied by Boehringer, Mannheim, Germany) : 2 μ mol/L umol/L pepstatin and 1 leupeptin, 2 phenylmethylsulphonyl fluoride (PMSF). Intact cells were lysed by passage through a French Pressure cell lb/in²). Cell debris was removed 000 centrifugation and lysates were diluted in column buffer to give a final concentration of protein/mL, prior to chromatography on a 2.6 cm x 20 cm column of amylose resin (New England Biolabs). The resin was washed with column buffer at 0.5 mL/min levels. The MBP-fused the returned until A_{280} recombinant proteins were eluted from the column by washing with column buffer containing 10 mmol/L 1maltose.

Fractions containing the recombinant proteins were pooled and then dialysed several times at 4° C against a low salt buffer (containing 25 mmol/L NaCl in 20 mmol/L TrisHCl, pH 8.0). The pooled fractions were then loaded at a flow rate of 0.5 mL/min onto a 1.6 x 10 cm anion exchange column (HP-Sepharose, Pharmacia, Sweden) connected Hi-Load to a system (Pharmacia). Proteins chromatography eluted from the column using a salt gradient (25 mmol/L to 500 mmol/L NaCl). Fractions giving high absorbance readings at A_{280} were exhaustively dialysed against distilled water at 4° C and analysed by SDS-PAGE.

Rabbit antisera :

Polyclonal rabbit antisera was prepared against total cell extracts of <u>H. pylori</u> strain 85P (Labigne et al., 1991) and <u>H. felis</u> (ATCC49179). Polyclonal rabbit antisera against recombinant protein preparations of <u>H. pylori</u> and <u>H. felis</u> urease subunits was produced by immunizing rabbits with 100 μ g of purified recombinant protein in Freund's complete adjuvant (Sigma). Four weeks later, rabbits were booster-immunized with 100 μ g protein in Freund's incomplete adjuvant. On week 6, the animals were terminally bled and the sera kept at -20° C.

Protein analyzes by SDS-PAGE and western blotting:

Solubilized cell extracts were analyzed on slab gels, comprising a 4.5% acrylamide stacking gel and a 10% resolving gel, according to the procedure of Laemmli. Electrophoresis was performed at 200 V on a mini-slab gel apparatus (Bio-Rad, USA).

Proteins were transferred to nitrocellulose paper in a Mini Trans-Blot transfer cell (Bio-Rad) set at 100 V for 1 h, with cooling. Nitrocellulose membranes were blocked with 5% (wt/vol) casein (BDH, England) in phosphate-buffered saline (PBS, pH 7.4) with gentle shaking at room temperature, for 2 h. Membranes were reacted at 4° C overnight with antisera diluted in 1% casein prepared in PBS. Immunoreactants were detected using specific biotinylated seondary antibodies and streptavidin-peroxidase conjugate (kirkegaard Parry Lab., Gaithersburg, USA). Reaction products were (Hyperfilm, autoradiographic film visualized on Amersham, France) using a chemiluminescence technique (ECL system, Amersham).

Protein concentrations were determined by the Bradford assay (Sigma Chemicals corp., St Louis, USA).

Animal experimentation:

Six week old female Swiss Specific Pathogen-Free (SPF) mice were obtained (Centre d'Elevage R. Janvier, Le-Genest-St-Isle, France) and maintained on a commercial pellet diet with water ad libitum. The intestines of the animals were screened for the absence of Helicobacter muridarum. For all orogastric administrations, 100 μ L aliquots were delivered to mice using 1.0 mL disposable syringes, to which polyethylene catheters (Biotrol, Paris, France) were attached.

Preparation of sonicated extracts and inocula from H. felis cultures:

H. felis bacteria were harvested in PBS and centrifuged at 5000 rpm, for 10 min in a Sorvall RC-5 centrifuge (Sorvall, USA) at 4° C. The pellets were washed twice and resuspended in PBS. Bacterial suspensions were sonicated as previously described and were subjected to at least one freeze-thaw cycle. Protein determinations were carried out on the sonicates.

To ensure a virulent culture of <u>H. felis</u> for protection studies, <u>H. felis</u> bacteria were maintained in vivo until required. Briefly, mice were inoculated three times (with 10¹⁰ bacteria/mL), over a period of 5 days. The bacteria were reisolated from stomach biopsies on blood agar medium (4 - 7 days' incubation in a microaerobic atmosphere at 37°C). Bacteria grown for two days on blood agar plates were harvested directly in peptone water (Difco, USA). Bacterial viability and motility was assessed by phase microscopy prior to administration to animals.

Mouse protection studies :

Fifty μg of recombinant antigen and 10 μg cholera holotoxin (Sigma Chemical Corp.), both resuspended in HCO_3 , were administrated orogastrically to mice on weeks 0, 1, 2 and 3. Mice immunized with sonicated \underline{H} . felis extracts (containing 400 - 800 μg of total protein) were also given 10 μg of cholera toxin. On week 5, half of the mice from each group were challenged with an inoculum of virulent \underline{H} . felis. The remainder of the mice received an additional "boost" immunization on week 15. On week 17 the latter were challenged with a culture of \underline{H} . felis.

Assessment of H. felis colonisation of the mouse:

Two weeks after receiving the challenge dose (ie. weeks 7 and 19, respectively) mice were sacrificed by spinal dislocation. The Stomachs were washed twice in sterile 0.8% NaCl and a portion of the gastric antrum from each stomach was placed on the surfaces of 12 cm x 12 cm agar plates containing a urea indicator medium (2% urea, 120 mg Na₂HPO₄, 80 mg KH₂PO₄, 1.2 mg phenol red, 1.5 g agar prepared in 100 mL). The remainder of each stomach was placed in formal-saline and stored until processed for histology. Longitudinal sections (4 μ m) of the stomachs were cut and routinely stained by the Giemsa technique. When necessary, sections were additionally stained by the Haematoxylin-Eosin and Warthin-Starry silver stain techniques;

The presence of <u>H. felis</u> bacteria in mouse gastric mucosa was assessed by the detection of urease activity (for up to 24 h) on the indicator medium, as well as by the screening of Giemsa-stained gastric sections that had been coded so as to eliminate observer bias. The numbers of bacteria in gastric sections were semi-quantitatively scored according to the following scheme: 0, no bacteria seen throughout

sections; 1, few bacteria (< 20) seen throughout; 2, occasional high power (H.P.) field with low numbers (< 20) of bacteria; 3, occasional H.P. field with low to moderate numbers (< 50) of bacteria; and 4, numerous (> 5) H.P. fields with high numbers of bacteria (> 50). Mononuclear cell infiltrates were scored as follows: 0, no significant infiltration; 1, infiltration of low numbers of mononuclear cells limited to the submucosa and muscularis mucosa; 2, infiltration of moderate numbers of mononuclear cells to the submucosa and muscularis mucosa, sometimes forming loose aggregates; and 3, infiltration of large numbers of mononuclear cells and featuring nodular agglomerations of cells.

RESULTS OF PART II EXPERIMENTS :

Expression of Helicobacter urease polypeptides in B. coli:

Fragments containing the sequences encoding the respective UreA gene products of \underline{H} . felis and \underline{H} . pylori were amplified by PCR and cloned in-phase with an ORF encoding the 42 kDa MBP, present on the expression vector pMAL. Sequencing of the PCR products revealed minor nucleotidic changes that did not, however, alter the deduced amino acid sequences of the respective gene products. E. coli MC1061 transformed with these recombinant plasmids (pILL919 and pILL920, respectively) expressed fusion proteins with predicted molecular weights of approximately 68 kDa. Following chromatography on affinity (amylose resin) and anion exchange gel media (Q-Sepharose), these proteins were purified to high degrees of purity (figure 1). The yield from 2-L cultures of recombinant E. coli cells was approximately 40 mg of purified antigen.

Similarly, the large UreB subunits of H. pylori and H. felis ureases were expressed in E. coli and pILL222, (plasmids pILL927 respectively) produced fusion proteins with predicted molecular weights of 103 kDa. The yield in these cases was appreciably lower than for the UreA preparations (approximately 20 mg was recovered from 2-L of bacterial culture). Moreover, problems associated with the cleavage of the UreB polypeptides from the MBP portion of the fusion proteins were encountered. These difficulties were attributed to the large sizes of the recombinant UreB polypeptides.

Analysis of the recombinant urease polypeptides:

Western blot analyses of the antigen preparations with rabbit polyclonal antisera raised to whole-extracts of <u>H. pylori</u> and <u>H. felis</u> bacteria demonstrated that the antigens retained immunogenicity to the homologous as well as heterologous antisera (figures 14 and 15). The antisera did not recognize the MBP component alone. Cross-reactivity between the urease polypeptides of <u>H. pylori</u> and <u>H. felis</u> was consistent with the high degrees of identity between the amino acid sequences of these proteins.

Rabbit polyclonal antisera raised against purified recombinant UreA and UreB proteins prepared from <u>H. pylori</u> and <u>H. felis</u> strongly reacted with the urease polypeptides present in whole-cell extracts of the bacteria (figure 16). As we had already observed, the UreB subunit of <u>H. felis</u> urease migrated slightly higher on SDS-PAGE gels than did that of <u>H. pylori</u> (figure 16).

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Preparation of H. felis inocula used in immunoprotection studies:

To ensure the virulence of <u>H. felis</u> bacterial inocula, bactera were reisolated from <u>H. felis</u>-infected mouse stomachs (see Materials and methods). The bacteria were passaged a minimum number of times in vitro. Stock cultures prepared from these bacteria, and stored at -80° C, were used to prepare fresh inocula for other mouse protection studies. This procedure ensured that the inocula used in successive experiments were reproducible.

Immunization of mice against gastric H. felisinfection:

Mice that had been immunized for three weeks with the given antigen preparations were divided into two lots and one half of these were challenged two weeks later with an <u>H. fellis</u> inoculum containing 10⁷ bacteria/mL. One group of animals that had been immunized with recombinant <u>H. felis</u> UreA were also challenged but, unlike the other animals, were not sacrificed until week 19.

a) Protection at week 5:

Eighty-five % of stomach biopsy samples from the control group of mice immunized with H. felis sonicate urease-negative preparations were and therefore have been protected from appeared to H. felis infection (table 4). This compared to 20% of those from the other control group of animals given MBP alone. The proportion of urease-negative stomachs for those groups of mice given the recombinant urease subunits varied from 70% (for H. pylori UreB) to 20% (for H. pylori UreA).

The levels of bacterial colonisation by H. felis also assessed from coded histological slides prepared from gastric tissue. Due to the striking helical morphology of H. felis bacteria, the organisms could be readily seen on the mucosal surfaces of both gastric pit and glandular regions of the stomach. Histological evidence indicated that the levels of protection in mice was lower than that observed by the biopsy urease test: 25% and 20% of gastric tissue felis immunized with H. mice from preparations of H. pylori UreB, respectively, were free of H. felis bacteria.

Amongst certain groups of these mice the preponderance of urease-negative biopsies, as well as lower histological scores for bacterial colonisation (unpublished data), suggested that an immunoprotective response had been elicited in the animals. This response, however, may have been insufficient to protect against the inoculum administered during the challenge procedure.

b) Protection at week 17:

The remaining mice, from each group of animals, were boosted on week 15. These mice were challenged at H. felis inoculum containing with an approximately 100-fold less bacteria than that used previously. Two weeks later all stomach biopsies from the MBP-immunized mice were urease-positive (table 4). In contrast, urease activity for gastric biopsies from mice immunized with the recombinant urease subunits varied from 50% for H. pylori UreA to 100% for H. felis UreB. The latter was comparable to the level of protection observed for the group of animals immunized sonicated extracts. Histological felis with evidence demonstrated that the UreB subunits of H.

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felis and <u>H. pylori</u> protected 60% and 25% of immunized animals, respectively. This compared with a level of 85% protection for mice immunized with <u>H. felis</u> sonicated extracts. Immunization of mice with recombinant <u>H. pylori</u> UreA did not protect the animals. Similarly, the stomachs of all <u>H. felis</u> UreA-immunized mice, that had been challenged at week 5, were heavily colonised with <u>H. felis</u> bacteria at week 19 (table 4).

The urease gastric biopsy test, when compared to histological analysis of gastric tissue sections, gave sensitivity and specificity values of 63% and 95%, respectively. Thus histology proved to be the more accurate predictor of <u>H. felis</u> infection in the mouse.

Cellular immune response in immunized stomachs:

In addition to the histological assessment of H. felis colonisation, mouse gastric tissue was also scored (from 0 to 3) for the presence of a mononuclear cell response. In mice immunized with MBP alone, a mild chronic gastritis was seen with small numbers of mononuclear cells restricted to the muscularis mucosa and to the submucosa of the gastric epithelium. contrast, there were considerable mononuclear cells present in the gastric mucosae from animals immunized with either the recombinant urease polypeptides, or with <u>H. felis</u> sonicate preparations. These inflammatory cells coalesced to form either loose aggregates, in the submucosal regions of the tissue, or nodular structures that extended into the regions of the gastric epithelia. mononuclear cell response did not appear to be related to the presence of bacteria as the gastric mucosae from the H. felis UreA-immunized mice, that were

heavily colonized with <u>H. felis</u> bacteria, contained little or no mononuclear cells.

Table ² The oligomeric primers used in PCR-based amplification of urease-encoding nucleotide sequences.

Prin	ner set	Nucleotide sequence (5' -> 3')
# 1	forw	CAU CCT* AAA ^G GAA ^G T ^C TA* GAT ^C AAA ^G T ^C TA* ATG
	rev	T ^C TC C ^T TT A*CG A*CG A*G ^C A ^T A ^{G,T} AT C ^T TT C ^T TT CAT CUA
#2	forw	CC GGA <u>GAA TTC</u> ATT AGC AGA AAA GAA TAT GTT TCT ATG <i>E</i> RT [¥]
	rev	AC GTT <u>CTG CAG</u> CTT ACG AAT AAC TTT TGT TGC TTG AGC
#3	forw	<u>GGA TCC</u> AAA AAG ATT TCA CG <i>Ba</i> mHI [¥]
	rev	GG <u>A AGC TT C TGC AG</u> G TGT GCT TCC CCA GTC HindIII [¥] Pstl [¥]

- Degenerated nucleotides in which all possible permutations of the genetic code were included (A, T, G, C).
- G,C,T The given nucleotides were degenerated with the specific base(s) shown.
 - [¥] Restriction sites introduced in the amplified fragments.

Table ³ Plasmids used

Plasmid	Vector	Relevant phenotype or character	Reference
pILL763	pILL570	9.5 kb fragment (Sau3a partial digest of H. pylori chromosome) (Sp ^R)	Cussac et al., 1991
pILL199	pILL575	35 kb fragment (Sau3A partial digest of H. felis chromosome)	Ferrero & Labigne,'93
pILL207	pILL570	11 kb fragment (Sau3A partial digest of pILL199)	This study
pILL919	pMAL-C2	0.8 kb BamHI-PstI a insert containing a nucleotide fragment encoding H. fe gene (ApR)	
pILL920	pMAL-C2	0.8 kb BamHI-PstI ^a insert containing PCR product encoding H. pylori ureA gene	This study
pILL927	pMAL-C2	1.8 kb EcoRI-PstIa PCR fragment encoding H. pylori ureB gene	This study
pILL213	pUC19	2 kb fragment resulting from Sau3A partial digest of pILL207 (Ap ^R)	This study
pILL219	pMAL-C2	1.4 kb DraI-HindIII ^b insert containing H. felis ureB (bases 657 - 1707)	This study
pILL 221	pMAL-C2	0.7 kb BamHI-PstI PCR fragment encoding H. felis ureB (bases 4 - 667)	This study
pILL222	pMAL-C2	1.35 kb PstI-PstI ^c fragment encoding H. felis ureB (bases 667 - 1707) from pILL219 cloned into linerized pILL221	

Table 4 Protection of mice by immunization with recombinant urease proteins.

Antigen		Protec	tion (%	%) a
	Ure	ase	Histo	logy
МВР	0 %	(0/10)	0 %	(0/10)
UreA H. pylori	50	(4/8)	0	(0/10)
UreA H. fclis b	12.5	(1/8)	0	(0/10)
UreB H. pylori	65	(5/8)	25	(2/8)
UreB H. felis	100	(7/7)	60	(5/7)
H. felis sonicate	100	(8/8)	85	(7/8)

- ^a Challenge inoculum dose was 10⁵ bacteria/mouse
- b Mice were challenged on week 5 (with 10⁷ bacteria) and were sacrificed on week 19.

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III- HELICOBACTER PYLORI hspA-B HEAT SHOCK GENE CLUSTER: NUCLEOTIDE SEQUENCE, EXPRESSION AND FUNCTION:

A homolog of the heat shock proteins (HSPs) of the GroEL class, reported to be closely associated with the urease of Helicobacter pylori (a nickel metalloenzyme), has recently been purified from H. pylori cells by Dunn et al, and Evans et al. (Infect. Immun. 60:1946, 1992, 1946 and 2125, respectively). Based on the reported N-terminal amino acid sequence of this immunodominant protein, degenerated oligonucleotides were synthesized in order to target the gene (hspB) encoding the GroEL-like protein in the chromosome of H. pylori strain 85P. Following gene amplification, a 108-base pair (bp)-fragment encoding the 36 first amino acids of the HspB protein was purified, and used a probe to identify in the H. pylori genomic bank a recombinant cosmid harboring the entire HspB encoding gene. The hspB gene was mapped to a 3.15 kilobases (kb) BglII restriction fragment of the pILL684 cosmid. The nucleotide sequence of that fragment subcloned into the pILL570 plasmid vector (pILL689) revealed the presence of two open reading frames (OFRs) designated hspA and hspB, the organization of which was very similar to be groESL bicistronic operons of other bacterial species. hspA and hspB encode polypeptides of 118 and 545 amino respectively, corresponding to calculated molecular masses of 13.0 and 58.2 kilodaltons (kDa), respectively. Amino acid sequence comparison studies revealed i) that the H. pylori HspA and HspB protein were highly similar to their bacterial homologs; ii) that the HspA H. pylori protein features a striking motif at the carboxyl terminus that other bacterial

GroEs-homologs lack; this unique motif consists of a series of eight histidine residues resembling metal binding domain, such a nickel binding. Surprisingly, immediately upstream of the gene cluster an insertion element was found that was absent in the H. pylori genome, and was positively selectionned during the cosmid cloning process. The IS5 was found to be involved in the expression of the hspA and hspB genes in pILL689. The expression of the HspA and HspB proteins from the pILL689 plasmid was analyzed in minicell-producing strain. Both polypeptides were shown to be constitutively expressed in the E. coli cells. When the pILL689 recombinant plasmid was introduced together with the H. pylori urease gene cluster into an E. coli host strain, an increase of urease activity was observed suggesting a close interaction between the heat shock proteins and the urease enzyme. Supporting the concept of a specific function for the HspA chaperone, was the fact that whereas a single hspB copy was found in the H. pylori genome, two copies of the hspA were found in the genome, one linked to the hspB gene and one unlinked to the hspB gene. Attempts to construct isogenic mutants of H. pylori in the hspA and the hspB gene were unsucesseful suggesting that these genes are essential for the survival of the bacteria.

EXPERIMENTAL PROCEDURES FOR PART III :

Bacterial strains, plasmids, and culture conditions:

The cloning experiments were performed with genomic DNA prepared from H. pylori strain 85P. H. pylori strain N6 was used as the recipient strain for the electroporation experiments because of its favorable transformability. E. coli strain HB101 or

strain MC1061 were used as a host for cosmid cloning and subcloning experiments, respectively. E. P678-54 was used for preparation of minicells. Vectors and recombinant plasmids used in this study are listed in Table 1. H. pylori strains were grown on horse blood agar plates, supplemented with vancomycin (10 mg/l), polymyxin B (2,500 U/I), trimethoprim (5 mg/l), and amphotericin B (4 mg/l). Plates were incubated at 37°C under microaerobic conditions in an anaerobic jar with a carbon dioxide generator envelope (BBL 70304). E. coli strains were grown in L-broth glucose (10 g of tryptone, 5 g of yeast extract, and 5 g of NaCI per liter; pH 7.0) or on L-agar plates (1.5 % agar) at 37°C. For measurement of urease activity, nitrogen-limiting medium used consisted the M9 minimal agar medium ammonium-free containing 0.4 % D-glucose as the carbon source, and freshly prepared filter-sterilized L-arginine added to 10 Antibiotic concentration of mM. concentrations for the selection of recombinant clones were as follows (in milligrams per liter) : kanamycin, 20 ; spectinomycin, 100 ; carbenicillin, 100.

Preparation of DNA:

Genomic DNA from H. pylori was prepared as previously described. Cosmid and plasmid DNAs were prepared by an alkaline lysis procedure followed by purification in cesium chloride-ethicium bromide gradients as previously described.

Cosmid cloning :

The construction of the cosmid gene bank of H. pylori 85P in E. coli HB101, which was used for the cloning of the H. pylori hspA-B gene cluster, has been described previously.

DNA analysis and cloning methodology:

Restriction endonucleases, T4 DNA ligase, I polymerase large (Klenow) fragment, and Taq polymerase were purchased from Amersham, T4DNA from Biolabs, polymerase and calf intestinal phosphatase from Pharmacia. All enzymes were used according to the instructions of the manufacturers. DNA fragments were separated on agarose gels run in Tris-acetate buffer. The 1-kb ladder from Bethesda Research Laboratories was used as a fragment size standard. When necessary, DNA fragments were isolated by electroelution from agarose gels as previously described and recovered from the migration buffer by means of an Elutip-d minicolumn (Schleicher and Schuell, Dassel, Germany). Basic DNA manipulations were performed according to the protocols described by Sambrook et al.

Hybridization:

Colony blots for screening of the H. pylori cosmid bank and for identification of subclones were prepared on nitrocellulose membranes (Schleicher and Schuell, Dassel, Germany) according to the protocol of Sambrook et al. (43). Radioactive labelling of PCRproducts was performed by random priming, using as primers the random hexamers from Pharmacia. Colony hybridizations were performed under high stringency conditions (5 x SSC, 0.1 % SDS, 50 % formamide, 42° C) (1 x SSC; 150 mM NaCl, 15 mM sodium citrate, pH 7.0). For Southern blot hybridizations, DNA fragments were transferred from agarose gels to nitrocellulose sheets $(0.45-\mu m \text{ pore size}$; Schleicher & Schuell, Inc.), and hybridized under low stringency conditions (5 x SSC, 0.1 % SDS, 30 or 40 % formamide, at 42° C with 32plabeled deoxyribonucleotide probes Hybridization was

revealed by autoradiography using Amersham Hyperfilm-MP.

DNA sequencing :

fragments of Appropriate plasmid DNA subcloned into M13 mp 18/19 vectors. Single stranded DNA was prepared by phage infection of E. coli strain JM101. Sequencing was performed by dideoxynucleotide chain termination method using the United States Biochemicals Sequenase kit. Both the M13 universal primer and additional specific primers (Fig.1) were used to sequence both the coding and non-coding DNA strands. Sequencing of double-stranded DNA was performed as previously described. Direct sequencing of PCR product was carried out following purification of the amplified, electroeluted product through an Elutip-d minicolumn (Schleicher & Schuell); The classical protocol for sequencing using the Sequenase kit was then used with the following modifications: PCR product was denatured by boiling annealing mixture containing 200 picomoles of the oligonucleotide used as primer and DMSO to the final concentration of 1 % for 3 minutes; the mixture was then immediatly cool on ice; the labeling step was performed in presence of manganese ions (mM).

Electroporation of H. pylori :

In the attempt to construct H. pylori mutants, plasmid constructions appropriate carrying targeted gene disrupted by a cassette containing a kanamycin resistance gene (aph3'-III), transformed into H. pylori strain N6 by means of previously electroporation as described. pSUS10 harboring the kanamycin disrupted flaA gene was used as positive control of electroporation. After

electroporation, bacteria were grown on non-selective plates for a period of 48 h in order to allow for the expression of the antibiotic resistance and then transferred onto kanamycin-containing plates. The selective plates were incubated for up to 6 days.

Polymerase chain reaction (PCR) :

PCRs were carried out using a Perkin-Elmer Cetus thermal cycler using the GeneAmp kit (Perkin-Elmer Cetus). Classical amplification reaction involved 50 picomoles (pmoles) of each primer and at least 5 pmoles of the target DNA. The target DNA was heat prior addition to the amplification denatured reaction. Reaction consisted of 25 cycles of the following three steps: denaturation (94° C for 1 minute), annealing (at temperatures ranging between 42 and 55° C, depending on the calculated melting temperatures of the primers, for 2 min), and extension (72° C for 2 min). When degenerated oligonucleotides were used in non stringent conditions, up to 1000 pmoles of each oligonucleotide were added, 50 cycles were carried out, and annealing was performed at 42° c.

Analysis of proteins expressed in minicells :

Minicells harboring the appropriate hybrid plasmid were isolated and labeled with [35 S] methionine (50 μ Ci/ml). Approximately 100,000 cpm of acetone-precipitable material was subjected to sodium dodecyl sulfate (SDS) -polyacrylamide gel electrophoresis in a 12.5 % gel. Standard proteins with molecular weights ranging from 94,000 to 14,000 (low< molecular-weights kit from Bio-Rad Laboratories) were run in parallel. The gel was stained and examined by fluorography, using En 3 Hance (New England Nuclear).

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Urease activity:

Urease activity was quantitated by the Berthelot reaction by using a modification of the procedure which has already been described. Urease activity was expressed as micromoles of urea hydrolyzed per minute per milligram of bacterial protein.

RESULTS OF PART III EXPERIMENTS :

Identification of a recombinant cosmid harboring the Helicobacter pylori GroEL-like heat shock protein encoding gene:

Based on the published N-terminal amino sequence of the purified heat shock protein of H. pylori, two degenerated oligonucleotides were synthesized target the gene of interest in the chromosome of H. pylori strain 85P. The first one 5' - G C N A A R G A RATHAARTTYTCNG-3' where N stands for the four nucleotides, R = A and G, Y = T and C, H = T, C, and A, is derived from for the first 8 amino acids of the protein (AKEIKFSD) ; the second one 5' - C R T T N C K N C C N C K N G G N C C C A T - 3', where K = G and T, corresponds to the complementary codons specifying the amino acid from position 29 to position 36 (MGPRGRNV, ref). The expected size for the PCR product was 108 base pairs (bp). The amplification reaction was performed under low stringency conditions as described in the "Materials and Methods" section, and led to the synthesis of six fragments with size ranging from 400 bp to 100 bp. The three smallest fragments were electroeluted from an acrylamide gel, and purified. Direct sequencing of the PCR products permitted the identification of a DNA encoding an amino acid sequence corresponding to the published sequence. This fragment was

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labeled and used as probe in colony hybridization to identify recombinant cosmids exhibiting homology to a 5' segment of the H. pylori GroEL-like encoding gene ; this gene was further designated hspB. The gene bank consists of 400 independent kanamycin-resistant coli transductants harboring recombinant cosmids. those one single clone hybridized with the probe, and harbored a recombinant plasmid designated pILL684, 46 kb in size. The low frequency observed when detecting the hspB gene (1 of 400) was unusual when compared that of several cloned genes which consistently detected in five to seven recombinant cosmids. In order to identify the hspB gene, fragments with sizes of 3 to 4 kb were generated by partial restriction of the pILL684 cosmid DNA with endonuclease Sau3A, purified, and ligated into the BglII site of plasmid vector pILL570. subclones, x were positive clones, and one was further studied (pILL689); it contains a 3.15 kb flanked by two BglII restriction sites, that was mapped in detail (Fig. 5). Using the PCR 32P labeled probe, the 5' end of the hspB gene was found to map to the 632 bp HindIII-SphI central restriction fragment of pILL689, indicating that one could expect the presence of the entire hspB gene in the pILL689 recombinant plasmid.

DNA sequence and deduced amino acid sequence of the H. pylori hspA-B gene cluster:

The 3200 bp of pILL689 depicted in Fig. 5 were sequenced by cloning into M13mp18 and M13mp19, the asymetric restriction fragments BglII-SphI, SphI-HindIII, HindIII-BglII; each cloned fragment was independently sequenced on both strands 16 oligonucleotide primers (Fig.1) were synthesized to

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confirm the reading and/or to generate sequences overlapping the independently sequenced fragments; these were used as primers in double-stranded-DNA sequencing analyses.

analysis of the sequence revealed distinct genetic elements. First the presence of two open reading frames (ORFs), depicted in figure 5, in transcribed the same direction, that designated hspA and hspB; The nucleotide sequence and the deduced amino acid sequence of the two ORFs are presented in Fig. 6. The first codon of hspA begins 323 bp upstream of the leftward HindIII site of pILL689 (Fig. 5) and is preceded by a Shine-Dalgarno ribosome-binding site (RBS) (GGAGAA). The hspA ORF codes for a polypeptide of 118 amino acids. The initiation codon for the hspB ORF nucleotides downstream the hspA stop codon ; it is preceded by a RBS site (AAGGA). The hspB ORF encodes a polypeptide of 545 amino acids and is terminated by a TAA codon followed by a palindromic resembling a rho-independent transcription terminator (free energy, $\Delta G = -19.8 \text{ kcal/mol}$) (Fig. 6). The Nterminal amino acid sequence of the deduced protein HspB was identical to the N-terminal sequence of the purified H.pylori heat shock protein previously the exception of published with the N-terminal methionine, which is absent from the purified protein and might be posttranslationally removed, resulting in a mature protein of 544 amino acids.

The deduced amino acid sequences of H. pylori HspA and HspB were compared to several amino acid sequences of HSPs of the GroES and GroEL class (Fig. 7). HspB exhibited high homology at the amino acid level with the Legionella pneumophila HtpB protein (82.9 % of similarities), with the Escherichia coli

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GroEL protein (81.0 % of similarities), with the Chlamydia psittaci or C. trachomatis HypB protein (79.4 % of similarities), with Clostridium perfringens Hsp60 protein (80.7 % of similarities), and to a the extent to GroEL-like proteins lesser of Mycobacterium. However, like almost all the GroEL homologs, H. pylori HspB demonstrated the conserved carboxyl-terminus glycine-methionine which was (MGGMGGMGGMM) recently shown to be dispensable in the E. coli GroEL chaperonin. degree of homology at the amino acid level between the pylori HspA protein and the other GroES-like proteins is shown in Fig. 7. The alignment shown features a striking motif at the carboxyl terminus of pylori HspA protein that other bacterial GroES-homologs lack. This unique highly charged motif consists of 27 additional amino acids capable of forming a loop between two double cystein residues; ot the 27 amino acids, 8 are histidine residues highly reminiscent of a metal binding domain.

second genetic element revealed sequence analysis, was the presence of an insertion sequence (IS5) 84 bp upstream of the hspA gene. The nucleotide sequence of this element matched perfectly that previously described for. IS5 in E. coli, with the 16 of a nucleotide (CTTGTTCGCACCTTCC) that corresponds to one of the two inverted repeats which flank the IS5 element. Because of the perfect match at the DNA level, we suspected that the IS5 was not initially present in the H. pylori chromosome, but had rather inserted upstream of the hspA-HspB gene cluster during the cloning process, a hypothesis that needed to be confirmed by further analyses.

Identification of the upstream sequence of the hspA-B gene cluster in H. pylori chromosome:

The presence of the IS5 was examined by gene amplification using two oligonucleotides, one being internal to the IS5 element and the other one downstream of the IS5 element (oligo #1 and #2, Fig. 6), to target a putative sequence i) in the chromosome of H. pylori strain 85P, ii) in the initial cosmid pILL684, and iii) in the 100 subclones resulting of Sau3A partial restriction of the recombinant cosmid. IS5 was absent from the chromosome of H. pylori, and was present in the very first subcultures of the E. coli strain harboring cosmid pILL684. Among the 100 pILL684 subclone derivatives which appeared to contain all or part of the IS5 sequence, we then looked for a subclone harboring the left end side of the IS5 plus the original upstream sequence of the hspA-hspB gene cluster. This screening was made by restriction analysis of the different Sau3A partial generated subclones. The restriction map of one (pILL694) of the plasmids fulfilling these criteria is shown in Fig. 5. The left end side of the IS5 nucleotide sequence was determined; the presence of a 4-bp duplication CTAA on both side of the 16-bp inverted repeats of the IS5 element (Fig. 6) allowed us to confirm the recent acquisition of the IS5 element by transposition. A 245-nucleotide sequence was then determined that mapped immediately upstream of the IS5 element (shown Fig. 6). This sequence consists of a non coding region in which the presence of a putative consensus heat shock promoter sequence was detected; it shows a perfectly conserved -35 region (TAACTCGCTTGAA) and a less consentaneous -10 region (CTCAATTA). Two oligonucleotides (#3 and #4, shown on Fig.2) were synthesized which mapped to

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sequences located on both side of the IS5 element the recombinant cosmid ; in these present oligonucleotides should lead to the amplification of a XXXXbp fragment when the IS5 sequence is present and a fragment in the absence of the IS5. The results of the PCR reaction using as target DNA the pILL684 cosmid, the pILL694 plasmid, and the H. pylori 85P chromosome fit the predictions (results not shown). Moreover, direct sequencing of the PCR product obtained from the H. pylori chromosome was performed and confirmed the upstream hspA-hspB reconstructed sequence shown further confirm (B). To the organization of the whole sequenced region, two probeswere prepared by gene amplification of the pILL689 plasmid using oligonucleotides #5 and #6, and #7 and #8 (Fig. 6).; they were used as probes in Southern hybridization experiments under low conditions against an HindIII digest of the H. pylori 85P chromosme. The results demonstrate that no other detectable rearrangement had occured during cloning process (data not shown). These experiments allowed us to demonstrate that whereas a single copy of the hspB gene was present in the chromosome of H. pylori strain 85, two copies of the hspA gene were detected by Southern hybridization.

Analysis of polypeptides expressed in minicells:

The pILL689 and the pILL692 recombinant plasmids and the respective cloning vectors pILL570, and pACYC177, were introduced by transformation into E. coli P678-54, a minicell-producing strain. The pILL689 and pILL692 plasmids (Fig. 5) contain the same 3.15-kb insert cloned into the two vectors. pILL570 contains upstream of the poly-cloning site a stop of transcription and of translation; the orientation of

the insert in pILL689, was made in such way that the transcriptinnal stop was located upstream of the IS5 fragment and therefore upstream of the hspA and HspB polypeptides Two that migrated polypeptides having apparent molecular weights of 60 kDa and 14 kDa were clearly detected in minicellexperiments from pILL689 and pILL692 (results not shown), whereas they were absent from the corresponding vectors ; these results indicated that the hspA and hspB genes were constitutively expressed promoter located within the IS5 were constitutively expressed from a promoter within the IS5 element. Moreover, whereas the amount of polypeptides visualized on the SDS gel was in good agreement with the copy number of the respective vectors, the intensity of the two polypeptidic bands suggested a polycistronic transcription of the two genes.

Attempts to understand the role of the Hspa and HspB proteins:

Two disruptions of genes were achieved in E. coli by inserting the Km cassette previously described within the hspA or the hspB gene of plasmids pILL686 and pILL691. This was done in order to return the disrupted genes in H. pylori by electroporation, and select for allelic replacement. The pILL696 resulting plasmid encoded a truncated form of the HspA protein, corresponding to the deletion of the Cterminal end amino acid sequence; in that plasmid the Km cassette was inserted in such way that the promoter of the Km gene could serve as promoter for the hspB downstream gene. The pILL687 and pILL688 plasmids resulted from the insertion of the Km cassette in either orientation within the hspB gene. None of these

constructs led to the isolation of kanamycin transformants of H. pylori strain N6, when purified pILL687, pILL688, pILL696 plasmids (Table 2, Fig. 5) were used in electroporation experiments, whereas the pSUS10 plasmid used as positive control always did. These results suggest the H. pylori HspA and HspB protein are essential proteins for the survival of H. pylori.

Because of i) the constant description in the literature of a close association of the HspB protein with the urease subunits ; -ii) the unique structure of the HspA protein with the C-terminal sequence reminiscent of a nickel binging domain, and iii) of the absence of viable hspA and/or hspB mutants of H. pylori, we attempted to demonstrate a role of the H. pylori Hsps proteins in relations with the H. pylori urease by functional complementation experiments in E. Plasmids pILL763 or pILL753 (both pILL570 derivatives, Table 5) encoding the urease gene cluster were introduced with the compatible pILL692 plasmid (pACYC177 derivative) that constitutively expresses HspA et HspB polypeptides as visualized minicells. In both complementations, the expression of the HspA and HspB proteins in the same E. coli cell allows to observe a three fold increase in the urease following induction of the urease genes on minimum medium supplemented with 10 mM L- Arginine as limiting nitrogen source.

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Table 5: Vectors and hybrid plasmids used in this study.

Plasmid	Vector	Size (kb)	Characteristics (a)	Origin or Reference
	p11.1.575	10	Mob, Cos, Km	
	p11.L570	5.3	Mob, Sp	
	pACYC177	3.9	Ap,Km	
p11.1.600	pBR322		Ap, Km, source of Km-cassette	
pH.1.684	pH.1.575		Mob, Kin, cosmid containing II. pylori hspA-B	
p11.1.685	pH.1.570		Mob, Sp, plasmid containing II. pylori hspB	San3A partial digest of pl1.1.684
p11.1.686	pUC19*c		Ap, plasmid containing 11. pylori lispB	
•	pUC19*(c)		Ap, Km, II. pylori lispit O Km-orientation A(b)	1.4-kb Smal-Smal plLL600 cloned into plLL686
p11.1.688	pUC19*(c)		Ap, Km, H. pylori hspB Ω Km- orientation B (b)	1.4-kb Smal-Smal pILL600 cloned into pILL686
p11.1.689	p1LL570		Mob, Sp, plasmid containing H. pylori hspA-B	Sau3A partial digest of pILL684
p11.1.691	pUC19**(c)		Ap, plasmid containing H.pylori lispA 1.3-kb	Sphl-Sphl plLL689 cloned into pUC19**
p11.1.692	pACYC177		Ap, Km, plasmid containing II. pylori hspA-B	3.15-kbBgIII pILL689 cloned into pACYC177
p11.1.694	p1LL570		Sp, plasmid containing left end of 1S5	Sau3A partial digest of p11.1.684
p11.1.696	pUC19**(c)		1p, Km, H. pylori lisp A O Km-orientation A (b)	1.4-kb Smal-Smal p1LL600 cloned into p1LL691
01SUS4	p1C20122		4p, Km, H. pylori fla A O Km	
p11.L753	p11.L570	16.5	Sp, plasmid containing ureA,B,C,D,E,F,G,H,I	•
p11.L763	p11.L570		Sp, plasmid containing ureA,B,E,F,G,II,I -	

(a) Mob, conjugative plasmid due to the presence of OriT; Ap, Km, and Sp, resistance to ampicillin, kanamycin, and spectinomycin, respectively; Cos, presence of lambda cos site.

(b) Orientation A indicates that the Kanamycin promoter initiates transcription in the same orientation as that of the gene where the

cassette has been inserted; orientation B, the opposite.
(c) pUC19* ane pUC19**; derivatives from pUC19 vector in which the the Sph1 and Hind111 site, respectively, have been end-filled by

using the Klenow polymerase and self religated.

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IV - EXPRESSION, PURIFICATION AND IMMUNOGENIC PROPERTIES OF H. PYLORI HSPA AND HSPB:

EXPERIMENTAL PROCEDURE FOR PART IV :

Expression and purification of recombinant fusion proteins:

The MalE-HspA, and MalE-HspB fusion proteins were expressed following the cloning of the two genes within the pMAL-c2 vector as described in "Results" section using the following primers : oligo #1 ccggagaattcAAGTTTCAACCATTAGGAGAAAGGGTC oligo #2 acgttctgcagTTTAGTGTTTTTTGTGATCATGACAGC oligo #3 ccqqaqaattcGCAAAAGAAATCAAATTTTCAGATAGC oligo #4 acgttctgcagATGATACCAAAAAGCAAGGGGGCTTAC Two liters of Luria medium containing glucose (30%) and ampicillin (100 μ g/ml) were inoculated with 20 ml of an overnight culture of strain MC1061 containing the fusion plasmid and incubated with shaking at 37°C. When the OD600 of the culture reached 0.5, IPTG (at a final concentration of 10 mM) was added, and the cells were incubated for a further 4 hours. Cells were harvested by centrifugation (5000 rpm for 30 min at 100 4°C), resuspended in ml of column buffer consisting of 10 mM Tris-HCl, 200 mM NaCl, 1 mM EDTA supplemented with protease inhibitors [(Leupeptin $(2\mu M)$ - Pepstatin $(2\mu m)$ - PMSF (1mM)- Aprotinin (1:1000 dilution)], and passed through a French press. After centrifugation (10,000 rpm for 20 min at 4°C), the supernatant were recovered and diluted (2-fold) with column buffer. The lysate was filtered through a 0.2 μm nitrocellulose filter prior to loading onto a preequilibrated amylose resin (22 x 2.5 cm). fusion proteins were eluted with a 10mM maltose solution prepared in column buffer, and the fractions

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containing the fusion proteins were pooled, dialyzed against distilled water, and lyophilized. Fusion proteins were resuspended in distilled water at a final concentration of 2 mg of lyophilized material/ml, and stored at -20°C. Concentration and purity of the preparations were controlled by the Bradford protein assay (Sigma Chemicals) and SDS-PAGE analyses.

Nickel binding properties of recombinant proteins:

E. coli MC1061 cells, containing either the pMAL-c2 vector or derivative recombinant plasmids, were grown in 100 ml-Luria broth in the presence of carbenicillin (100 μ g/ml). The expression of the genes was induced with IPTG for four hours. The cells were centrifuged and the pellet was resuspended in 2 ml of Buffer A (6M guanidine hydrochloride, 0.1 M NaH2PO4, 0.01Tris, pH8.0). After gentle stirring for one hour at room temperature, the suspensions were centrifuged at 10,000 g for 15 min at 4°C. A 1.6 ml aliquot of Nickel-Nitrilo-Tri-Acetic resin (Nickel-NTA, QIA express), previously equilibrated in Buffer A, added to the supernatant and this mixture was stirred at room temperature for one hour prior to loading onto a column. The column was washed with 20 ml buffer A, then 30 ml buffer B (8M urea, 0.1M Na-phosphate, 0.01MTris-HCl, pH8.0). The proteins were successively with the same buffer as buffer B adjusted to pH 6.3 (Buffer C), pH 5.9 (Buffer D) and pH 4.5 (Buffer E) and Buffer F (6M guanidine hydrochloride, 02M acetic acid). Fifty μ l of each fraction were mixed with 50 µl of SDS buffer and loaded on SDS gels.

Human sera :

Serum samples were obtained from 40 individuals, 28 were <u>H. pylori</u>-infected patients as confirmed by a positive culture for <u>H. pylori</u> and histological examination of the biopsy, and 12 were uninfected patients. The sera were kindly provided by R. J. Adamek (University of Bochum, Germany).

Immunoblotting:

Upon completion of SDS-PAGE runs in a Minielectrophoresis cell, proteins transferred to nitrocellulose paper in a Mini Trans-Blot transfer cell (Bio-Rad) set at 100 V for 1 h (with cooling). Immunostaining was performed as. previously described (Ferrero et al., 1992), except ECL Western blotting detection (Amersham) was used to visualize reaction products . Human sera and the rabbit antiserum, raised against a whole-cell extract of H. pylori strain 85P, were diluted 1:1000 and 1:5000, respectively, in 1% (w/v)casein prepared in phosphate-buffered saline (PBS, pH7.4).

Serological methods [enzyme-linked immunosorbent assay, (ELISA)]:

The following quantities of antigens were absorbed onto 96-well plates (Falcon 3072): 2.5 μ g of protein MalE, 5 μ g of MalE-HspA, or 2.5 μ g of MalE-HspB. The plates were left overnight at 4°C, then washed 3 times with ELISA wash solution (EWS) [1% PBS containing 0.05% (v/v) Tween 20]. Saturation was achieved by incubating the plates for 90 min at 37°C in EWS supplemented with 1% milk powder. Wells were again washed 3 times with EWS and then gently agitated for 90 min at 37°C in the presence of human sera (diluted 1:500 in EWS with 0.5% milk powder), under

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agitation. Bound imunoglobulins were detected by incubation for 90 min at 37°C with biotinylated secondary antibody (goat anti-human IgG, IgA or IgM diluted [1:1000] in EWS supplemented with 0.5% milk powder) in combination with streptavidin-peroxidase (1:500) (Kirkegaard and Perry Lab.). Bound peroxidase was detected by reaction with the citrate substrate and hydrogen peroxide. Plates were incubated in the dark, at room temperature, and the optical density at 492 nm was read at intervals of 5, 15 and 30 min in an ELISA plate reader. After 30 min, the reaction was stopped by the addition of hydrochloric acid to a final concentration of 0.5M.

RESULTS OF PART IV EXPERIMENTS:

Construction of recombinant plasmids producing inducible MalE-HspA, and HspB fusion proteins:

The oligonucleotides #1 and #2 (hspA) and #3 and #4 (hspB) were used to amplify by PCR the entire hspA and the hspB genes, respectively. The PCR products were electroeluted, purified and restricted with EcoRI and PstI. The restricted fragments (360 bp and 1600 bp in size, respectively) were then ligated into the EcoRI-PstI restricted pMAL-c2 vector to plasmids designated pILL933 and pILL934, respectively. Following induction with IPTG, and purification of the soluble protein on amylose columns, fusion proteins of the expected size (55 kDa for pILL933 [figure 17], and 100 kDa for pILL9334) were visualized on SDS-PAGE gels. Each of these corresponded to the fusion of the MalE protein (42.7 kDa) with the second amino-acid of each of the Hsp polypeptides. The yield expression of the fusion proteins was 100 mg

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MalE-HspA and 20 mg for MalE-HspB when prepared from 2 liters of broth culture.

Study of the antiquenicity of the HspA and HspB fusion proteins, and of the immunogenicity of HspA and HspB in patients infected with H. pylori:

In order to determine whether the fusion proteins were still antigenic, each was analyzed by Western blot with rabbit antiserum raised against the MalE protein and a whole-cell extract of <u>H. pylori</u> strain 85P. Both fusion proteins were immunoreactive with antibody to MalE (not shown) and with the anti-<u>H. pylori</u> antiserum. The anti-<u>H. pylori</u> antiserum did not recognize the purified MalE protein (figure 18). These results demonstrated that the fusion proteins retained their antigenic properties; in addition, whereas the HspB protein was known to be immunogenic, this is the first demonstration that HspA <u>per se</u> is immunogenic in rabbits.

In the same way, in order to determine whether the HspA and HspB polypeptides were immunogenic in humans, the humoral immune response against HspA and/or HspB in patients infected with H. pylori was analyzed and compared to that of uninfected persons using Western immunoblotting assays and enzyme-linked immunosorbent assays (ELISA). None of the 12 sera of H. __pylori-negative persons gave a immunoblot signal with MalE, MalE-HspA, or MalE-HspB proteins (figure 18). In contrast, of 28 sera from H. pylori-positive patients, 12 (42.8%) reacted with the HspA protein whilst 20 (71.4%) recognized the HspB protein. All of the sera that recognized HspA also reacted with the HspB protein. No association was observed between the immune response and the clinical presentation of the H. pylori infection although such

a conclusion might be premature because of the small number of strains analyzed.

Nickel binding properties of the fused MalE-HspA protein:

MBP-HspA recombinant protein expressed following induction with IPTG, was purified from a whole cell extract by one step purification on nickel affinity column whereas the MBP alone, nor MBP-HspB exhibited this property. Figure 18 illustrates the one step purification of the MBP-HspA protein that was eluted as a monomer at pH6.3, and as a monomer at pH4.5. The unique band seen in panel 7 and the two bands seen in panel 5 were both specifically recognized with anti-HspA rabbit sera. This suggested that the nickel binding property of the fused MBP-HspA protein might be attributed to the C-terminal sequence os HspA which is rich in Histidine and Cysteine residues.

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SEQUENCE LISTING

- (1) GENERAL INFORMATION:
 - (i) APPLICANT:
 - (A) NAME: INSTITUT PASTEUR
 - (B) STREET: 25-28 rue du Dr Roux
 - (C) CITY: PARIS CEDEX 15
 - (E) COUNTRY: FRANCE
 - (F) POSTAL CODE (ZIP): 75724
 - (G) TELEPHONE: 45.68.80.94
 - (H) TELEFAX: 40.61.30.17
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 - (B) STREET: 101 rue de Tolbiac
 - (C) CITY: PARIS CEDEX 13
 - (E) COUNTRY: FRANCE
 - (F) POSTAL CODE (ZIP): 75654
 - (G) TELEPHONE: 44.23.60.00
 - (H) TELEFAX: 45.85.07.66
 - (ii) TITLE OF INVENTION: IMMUNOGENIC COMPOSITIONS AGAINST HELICOBACTER INFECTION, POLYPEPTIDES FOR USE IN THE COMPOSITIONS AND NUCLEIC ACID SEQUENCES ENCODING SAID POLYPEPTIDES.
 - (iii) NUMBER OF SEQUENCES: 8
 - (iv) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25 (EPO)
 - (v) CURRENT APPLICATION DATA:
 APPLICATION NUMBER: EP 93401309.5
- (2) INFORMATION FOR SEQ ID NO: 1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2619 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (ix) FEATURE:
 - (A) NAME/KEY: misc_feature
 - (B) LOCATION: 31..36
 - (D) OTHER INFORMATION: /standard_name= "Shine-Dalgarno sequence"

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	(ix	(A) N B) L	AME/ OCAT	KEY: ION:	43.	. 753		tand	ard_	name	.= "U	IRE A	, a		
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	(xi) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0: 1	:					
TGA	TAGC	TTG	GCTA	CCAA	TA G	AAAT	TCAA	T AA	GGAG	TTTA	GG			CTA Leu		54
CCT Pro 5	Lys	GÀA Glu	CTA Leu	GAC Asp	AAG Lys 10	Leu	ATG Met	CTC Leu	CAT His	TAT Tyr 15	Ala	GGC Gly	AGA Arg	TTG Leu	GCA Ala 20	102
GAA Glu	GAA Glu	CGC Arg	TTG Leu	GCG Ala 25	CGT Arg	GGT Gly	GTG Val	AAA Lys	CTC Leu 30	AAT Asn	TAC	ACC Thr	GAA Glu	GCG Ala 35	GTC Val	150
GCG Ala	CTC Leu	ATT Ile	AGC Ser 40	GGG Gly	CGT Arg	GTG Val	ATG Met	GAA Glu 45	AAG Lys	GCG Ala	CGT Arg	GAT Asp	GGT Gly 50	AAT Asn	AAA Lys	198
AGC Ser	GTG Val	GCG Ala 55	GAT Asp	TTG Leu	ATG Met	CAA Gln	GAA Glu 60	Gly	Arg	Thr	TGG Trp	Leu	Lys	AAA Lys	GAA Glu	246
AAT Asn	GTG Val 70	ATG Met	GAC Asp	GGC Gly	GTA Val	GCA Ala 75	AGC Ser	ATG Met	ATT Ile	CAT His	GAA Glu 80	GTG Val	GGG Gly	ATT Ile	GAA Glu	294
GCT Ala 85	AAC Asn	TTC Phe	CCC Pro	GAT Asp	GGA Gly 90	ACC Thr	AAG Lys	CTT Leu	GTA Val	ACT Thr 95	ATC Ile	CAC His	ACT Thr	CCG Pro	GTA Val 100	342
GAG Glu	GAT Asp	AAT Asn	GGC Gly	AAA Lys 105	TTA Leu	GCC Ala	CCC Pro	GGC Gly	GAG Glu 110	GTC Val	TTC Phe	TTA Leu	AAA Lys	AAT Asn 115	GAG Glu	390
GAC Asp	ATT Ile	ACT Thr	ATT Ile 120	AAC Asn	GCC Ala	GGC Gly	AAA Lys	GAA Glu 125	GCC Ala	ATT Ile	AGC Ser	TTG Leu	AAA Lys 130	GTG Val	AAA Lys	438

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	AAA Lys															486
	GTG Val 150															534
	CTA Leu															582
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	TTT Phe															678
	GGC Gly															726
	TGT Cys 230									GAA/	AAA (_	AA AA 7s Ly		774
ATT Ile	TCA Ser 5	CGA Arg	AAA Lys	GAA Glu	TAT Tyr	GTT Val 10	TCT Ser	ATG Met	TAT Tyr	GGT Gly	CCC Pro 15	ACT Thr	ACC Thr	GGG Gly	GAT Asp	822
	GTT Val															870
	ACC Thr															918
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	AAT Asn															1110

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1								GGC Gly		1350
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I								CAC His		1590
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								CAC His		1734
								AGG Arg		1782

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											86					
CG(Arg 34(grr	C CA	A ACT	T ATO	345	a Ala	GAA Glu	GA(As _I	C CAA	CTO Let 350	ı His	GAC S Asp	C ATO	GGG Gl	G ATC y Ile 355	1830.
TT1 Phe	TC: Sei	T ATO	C ACC	Ser 360	Ser	GAC Asp	TCT Ser	CAC Glr	G GCT n Ala 365	Met	G GGA	A CGC	GTA Val	GG(G13	GAG Glu	1878
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GTG Val	ACC Thr	GCA Ala	CAT His 535	ATT Ile	GAT Asp	GTC Val	Asn	CCT Pro 540	GAA Glu	ACC Thr	TAT Tyr	Lys	GTG Val 545	AAA Lys	GTG Val	2406
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		Tyr			TTC Phe	TAG		CTA	AGGA	GGGG	GA T	AGAG	GGGG	Т		2502
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TTT	TTTC	GTG	TTTT	ATAC	CG C	GTTG	AAAC	C CT	CAAA	тстт	TAC	CAAA	AGG .	ATGG	TAA	2619
(2)	INF	ORMA	TION	FOR	SEQ	ID !	NO:	2:							·	
		(, (,	A) L B) T	ENGT YPE :	CHAI H: 2: amii OGY:	37 au no a	mino cid									
	(ii) MO	LECU	LE T	YPE:	pro	tein									
	(vi				OURCI		icoba	acte:	r fe	lis						
	(xi) SE	QUEN	CE D	ESCR:	IPTI(ON:	SEQ :	ID N	0: 2	:					
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Thr	Glu	Ala 35	Val	Ala	Leu	Ile	Ser 40	Gly	Arg	Val	Met	Glu 45	Lys	Ala	Arg	
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His	Thr	Pro	Val 100	Glu	Asp	Asn	Gly	Lys 105	Leu	Ala	Pro	Gly	Glu 110	Val	Phe	
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Ser	Phe	Cys	Lys	Arg 165	Leu	Asp	Ile	Ala	Ser 170	Gly	Thr	Ala	Val	Arg 175	Phe	

Glu Pro Gly Glu Glu Lys Ser Val Glu Leu Ile Asp Ile Gly Gly Asn 180 185 190

Lys Arg Ile Tyr Gly Phe Asn Ser Leu Val Asp Arg Gln Ala Asp Ala 195 200 205

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- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 569 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (vi) ORIGINAL SOURCE :
 - (A) ORGANISM: Helicobacter felis
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:

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Glu His Asp Cys Thr Thr Tyr Gly Glu Glu Ile Lys Phe Gly Gly 35 40 45

Lys Thr Ile Arg Asp Gly Met Ser Gln Thr Asn Ser Pro Ser Ser Tyr 50 60 .

Glu Leu Asp Leu Val Leu Thr Asn Ala Leu Île Val Asp Tyr Thr Gly
65 70 75 80

Ile Tyr Lys Ala Asp Ile Gly Ile Lys Asp Gly Lys Ile Ala Gly Ile 85 90 95

Gly Lys Ala Gly Asn Lys Asp Met Gln Asp Gly Val Asp Asn Asn Leu 100 105 110

Cys Val Gly Pro Ala Thr Glu Ala Leu Ala Ala Glu Gly Leu Ile Val

Thr Ala Gly Gly Ile Asp Thr His Ile His Phe Ile Ser Pro Gln Gln 130 135 140

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455

90 Ala Ser Ile Pro Thr Pro Gln Pro Val Tyr Tyr Arg Glu Met Phe Gly 465 His His Gly Lys Asn Lys Phe Asp Thr Asn Ile Thr Phe Val Ser Gln 485 490 Ala Ala Tyr Lys Ala Gly Ile Lys Glu Glu Leu Gly Leu Asp Arg Ala 500 Ala Pro Pro Val Lys Asn Cys Arg Asn Ile Thr Lys Lys Asp Leu Lys 520 Phe Asn Asp Val Thr Ala His Ile Asp Val Asn Pro Glu Thr Tyr Lys 540 Val Lys Val Asp Gly Lys Glu Val Thr Ser Lys Ala Ala Asp Glu Leu 545 550 555 Ser Leu Ala Gln Leu Tyr Asn Leu Phe 565 (2) INFORMATION FOR SEQ ID NO: 4: (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 2284 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 124..477
 - (D) OTHER INFORMATION: /standard_name=."H. pylori Hsp A"
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 - (A) NAME/KEY: CDS

20

- (B) LOCATION: 506..2143
- (D) OTHER INFORMATION: /standard_name= "H. pylori Hsp B"
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TGT	CGCT	CAA (GAATA	ACTA	G CC	GCTA/	\ATT	r cta	ATTT	TTAT	TAT	CAAA	ACT :	[AGG	AGAACT	120
GAA			TTT Phe													168
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CAT His	ACA Thr	GGT Gly	AAT Asn	CAT His 100	GAT Asp	CAT His	AAA Lys	CAT His	GCT Ala 105	AAA Lys	GAG Glu	CAT His	GAA Glu	GCT Ala 110	Cys	456
TGT Cys	CAT His	GAT Asp	CAC His 115	AAA Lys	AAA Lys	CAC	TAA	AAA	CAT 1	[ATT	ATTA	AG G	ATAC		ATG Met 1	508
GCA Ala	AAA Lys	GAA Glu	ATC Ile 5	AAA Lys	TTT Phe	TCA Ser	GAT Asp	AGC Ser 10	GCA Ala	AGA Arg	AAC Asn	CTT Leu	TTA Leu 15	TTT Phe	GAA Glu	556
GGC Gly	GTA Val	AGA Arg 20	CAA Gln	CTC Leu	CAT His	GAC Asp	GCT Ala 25	GTC Val	AAA Lys	GTA Val	ACC Thr	ATG Met 30	GGG Gly	CCA Pro	AGA Arg	604
GGC Gly	AGG Arg 35	AAC Asn	GTG Val	TTG Leu	ATC Ile	CAA Gln 40	AAA Lys	AGC Ser	TAT Tyr	GGC Gly	GCT Ala 45	CCA Pro	AGC Ser	ATC Ile	ACC Thr	652
AAA Lys 50	GAC Asp	GGC Gly	GTG Val	AGC Ser	GTG Val 55	GCT Ala	AAA Lys	GAG Glu	ATT Ile	GAA Glu 60	TTA Leu	AGT Ser	TGC Cys	CCC Pro	GTG Val 65	700
GCT Ala	AAC Asn	ATG Met	GGC Gly	GCT Ala 70	CAG Gln	CTC Leu	GTT Val	AAA Lys	GAA Glu 75	GAT Asp	GCG Ala	AGC Ser	AAA Lys	ACC Thr 80	GCT Ala	748
GAT Asp	GCC Ala	GCC Ala	GGC Gly 85	GAT Asp	GGC Gly	ACG Thr	ACC Thr	ACA Thr 90	GCG Ala	ACC Thr	GTG Val	CTG Leu	GCT Ala 95	TAT Tyr	AGC Ser	796
ATT Ile	TTT Phe	AAA Lys 100	GAG Glu	GGC Gly	TTG Leu	AGG Arg	AAT Asn 105	ATC Ile	ACG Thr	GCT Ala	GGG Gly	GCT Ala 110	AAC Asn	CCT Pro	ATT Ile	844
GAA Glu	GTG Val	AAA Lys	CGA Arg	GGC Gly	ATG Met	GAT Asp	AAA Lys	GCG Ala	CCT Pro	GAA Glu	GCG Ala	ATC Ile	ATT Ile	AAT Asn	GAG Glu	892

											12					
CTT Leu 130	Lys	A AAA S Lys	A GCC s Ala	G AGO	Lys 135	Lys	GTC Val	GGC Gly	GGT Gly	C AAA 7 Lys 140	Gli	A GAA	ATO	C ACC	CAA Gln 145	940
GTA Val	GCC Ala	ACC Thr	ATI	Ser 150	' Ala	AAC Asn	Ser	GAI Asp	CAC His 155	: Asr	ATO	GGG Gly	AAA Lys	A CT(5 Leu 16(C ATC L Ile	988
GCT Ala	GAC Asp	GCT Ala	ATG Met 165	GLu	AAA Lys	GTG Val	GGT Gly	AAA Lys 170	Asp	GGC Gly	GTC Val	ATC	ACC Thr 175	Val	GAA Glu	1036
GAA Glu	GCT Ala	Lys 180	GIA	ATT Ile	GAA Glu	GAT Asp	GAA Glu 185	Leu	GAT Asp	GTC Val	GTA Val	GAA Glu 190	Gly	ATG Met	CAA Gln	1084
TTT Phe	GAT Asp 195	Arg	GGC Gly	TAC Tyr	CTC Leu	TCC Ser 200	CCT Pro	TAC Tyr	TTT Phe	GTA Val	ACC Thr 205	Asn	GCT Ala	GAG Glu	AAA Lys	1132
ATG Met 210	ACC Thr	GCT Ala	CAA Gln	TTG Leu	GAT Asp 215	AAC Asn	GCT Ala	TAC Tyr	ATC Ile	CTT Leu 220	TTA Leu	ACG Thr	GAT Asp	AAA Lys	AAA Lys 225	1180
ATC Ile	TCT Ser	AGC Ser	ATG Met	AAA Lys 230	GAC Asp	ATT Ile	CTC Leu	CCG Pro	CTA Leu 235	CTA Leu	GAA Glu	AAA Lys	ACC Thr	ATG Met 240	AAA Lys	1228
GAG Glu	GGC Gly	AAA Lys	CCG Pro 245	CTT Leu	TTA Leu	ATC Ile	ATC Ile	GCT Ala 250	GAA Glu	GAC Asp	ATT Ile	GAG Glu	GGC Gly 255	GAA Glu	GCT Ala	1276
TTA Leu	ACG Thr	ACT Thr 260	CTA Leu	GTG Val	GTG Val	AAT Asn	AAA Lys 265	TTA Leu	AGA Arg	GGC Gly	GTG Val	TTG Leu 270	AAT Asn	ATC Ile	GCA Ala	1324
A14	GTT Val 275	AAA Lys	GCT Ala	CCA Pro	GGC Gly	TTT Phe 280	GGG Gly	GAC Asp	AGG Arg	AGA Arg	AAA Lys 285	GAA Glu	ATG Met	CTC Leu	AAA Lys	1372
GAC Asp 290	ATC Ile	GCT Ala	GTT Val	TTA Leu	ACC Thr 295	GGC Gly	GGT Gly	CAA Gln	GTC Val	ATT Ile 300	AGC Ser	GAA Glu	GAA Glu	TTG Leu	GGC Gly 305	1420
TTG . Leu	AGT Ser	CTA Leu	GAA Glu	AAC Asn 310	GCT Ala	GAA Glu	GTG Val	GAG Glu	TTT Phe 315	TTA Leu	GGC Gly	AAA Lys	GCG Ala	AAG Lys 320	ATT Ile	1468
CTC / Val	ATT Ile	GAC Asp	AAA Lys 325	GAC Asp	AAC Asn	ACC Thr	Thr	ATC Ile 330	GTA Val	GAT Asp	GGC Gly	Lys	GGC Gly 335	CAT His	AGC Ser	1516
CAT (Asp	GTC Val 340	AAA Lys	GAC Asp	AGA Arg	Val .	GCG Ala 345	CAA Gln	ATC Il	AAA Lys	ACC Thr	CAA Gln 350	ATT Ile	GCA Ala	AGC Ser	1564

											93					
		A AGO r Ser														1612
	u Se	r GGC r Gly														1660
		G AAA t Lys														1708
		G GCC a Ala		Glu												1756
		C GCC g Ala 420	Ala													1804
		C TAT y Tyr 5														1852
	e Al	T ATO														1900
		A CA(s His			His											1948
		C ATO		Lys												1996
		T TTA a Let 500	ı Gln					Val								2044
		C ACC a Thi					Lys									2092
	o As	T ATO				Gly										2140
TA	AGCC	CCCT	TGCI	TTTI	GG T	ATCA	TCTG	C TT	TTAA	AATC	CAT	CTTC	TAG .	AATC	CCCCCT	2200
TC	TAAA	ATCC	CTTI	TTTG	GG G	GGTG	CTTT	T GG	TTTG	ATAA	AAC	CGCT	ccc ·	TTTT	AAAAAC	2260
GC	GCAA	CAAA	AAAC	TCTG	TT A	AGC										2284

(2) INFORMATION FOR SEQ ID NO: 5:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 545 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM : H. pylori
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:
- Met Ala Lys Glu Ile Lys Phe Ser Asp Ser Ala Arg Asn Leu Leu Phe 1 5 10 15
- Glu Gly Val Arg Gln Leu His Asp Ala Val Lys Val Thr Met Gly Pro 20 25 30
- Arg Gly Arg Asn Val Leu Ile Gln Lys Ser Tyr Gly Ala Pro Ser Ile 35 40 45
- Thr Lys Asp Gly Val Ser Val Ala Lys Glu Ile Glu Leu Ser Cys Pro
 50 55 60
- Val Ala Asn Met Gly Ala Gln Leu Val Lys Glu Asp Ala Ser Lys Thr
 65 70 75 80
- Ala Asp Ala Ala Gly Asp Gly Thr Thr Thr Ala Thr Val Leu Ala Tyr
 85 90 95
- Ser Ile Phe Lys Glu Gly Leu Arg Asn Ile Thr Ala Gly Ala Asn Pro 100 105 110
- Ile Glu Val Lys Arg Gly Met Asp Lys Ala Pro Glu Ala Ile Ile Asn 115 120 125
- Glu Leu Lys Lys Ala Ser Lys Lys Val Gly Gly Lys Glu Glu Ile Thr 130 135 140
- Gln Val Ala Thr Ile Ser Ala Asn Ser Asp His Asn Ile Gly Lys Leu 145 150 155 160
- Ile Ala Asp Ala Met Glu Lys Val Gly Lys Asp Gly Val Ile Thr Val 165 170 175
- Glu Glu Ala Lys Gly Ile Glu Asp Glu Leu Asp Val Val Glu Gly Met 180 185 190
- Gln Phe Asp Arg Gly Tyr Leu Ser Pro Tyr Phe Val Thr Asn Ala Glu 195 200 205
- Lys Met Thr Ala Gln Leu Asp Asn Ala Tyr Ile Leu Leu Thr Asp Lys 210 215 220

- 1

- Lys Ile Ser Ser Met Lys Asp Ile Leu Pro Leu Leu Glu Lys Thr Met 225 230 235 240
- Lys Glu Gly Lys Pro Leu Leu Ile Ile Ala Glu Asp Ile Glu Gly Glu 245 250 255
- Ala Leu Thr Thr Leu Val Val Asn Lys Leu Arg Gly Val Leu Asn Ile 260 270
- Ala Ala Val Lys Ala Pro Gly Phe Gly Asp Arg Arg Lys Glu Met Leu 275 280 285
- Lys Asp Ile Ala Val Leu Thr Gly Gly Gln Val Ile Ser Glu Glu Leu 290 295 300
- Gly Leu Ser Leu Glu Asn Ala Glu Val Glu Phe Leu Gly Lys Ala Lys 305 310 315 320
- Ile Val Ile Asp Lys Asp Asn Thr Thr Ile Val Asp Gly Lys Gly His 325 330 335
- Ser His Asp Val Lys Asp Arg Val Ala Gln Ile Lys Thr Gln Ile Ala 340 345 350
- Ser Thr Thr Ser Asp Tyr Asp Lys Glu Lys Leu Gln Glu Arg Leu Ala 355 360 365
- Lys Leu Ser Gly Gly Val Ala Val Ile Lys Val Gly Ala Ala Ser Glu 370 375 380
- Val Glu Met Lys Glu Lys Lys Asp Arg Val Asp Asp Ala Leu Ser Ala 385 390 395 400
- Thr Lys Ala Ala Val Glu Glu Gly Ile Val Ile Gly Gly Gly Ala Ala
 405 410 415
- Leu Ile Arg Ala Ala Gln Lys Val His Leu Asn Leu His Asp Asp Glu 420 425 430
- Lys Val Gly Tyr Glu Ile Ile Met Arg Ala Ile Lys Ala Pro Leu Ala
 435
 440
 445
- Gln Ile Ala Ile Asn Ala Gly Tyr Asp Gly Gly Val Val Val Asn Glu 450 455 460
- Val Glu Lys His Glu Gly His Phe Gly Phe Asn Ala Ser Asn Gly Lys 465 470 475 480
- Tyr Val Asp Met Phe Lys Glu Gly Ile Ile Asp Pro Leu Lys Val Glu 485 490 495
- Arg Ile Ala Leu Gln Asn Ala Val Ser Val Ser Ser Leu Leu Thr 500 505 510
- Thr Glu Ala Thr Val His Glu Ile Lys Glu Glu Lys Ala Ala Pro Ala 515 520 525

M t Pro Asp Met Gly Gly Met Gly Gly Met Gly Gly Met Gly Gly Met 530 535 . 540

Met 545

- (2) INFORMATION FOR SEQ ID NO: 6:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 118 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM : H. pylori
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:

Met Lys Phe Gln Pro Leu Gly Glu Arg Val Leu Val Glu Arg Leu Glu
1 5 10 15

Glu Glu Asn Lys Thr Ser Ser Gly Ile Ile Ile Pro Asp Asn Ala Lys
20 25 30

Glu Lys Pro Leu Met Gly Val Val Lys Ala Val Ser His Lys Ile Ser 35 40 45

Glu Gly Cys Lys Cys Val Lys Glu Gly Asp Val Ile Ala Phe Gly Lys 50 55 60

Tyr Lys Gly Ala Glu Ile Val Leu Asp Gly Val Glu Tyr Met Val Leu
65 70 75 80

Glu Leu Glu Asp Ile Leu Gly Ile Val Gly Ser Gly Ser Cys Cys His
85 90 95

Thr Gly Asn His Asp His Lys His Ala Lys Glu His Glu Ala Cys Cys
100 105 110

His Asp His Lys Lys His
115

- (2) INFORMATION FOR SEQ ID NO: 7:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 591 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)